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APPENDIX A

GAS-PHASE ATMOSPHERIC CHEMISTRY OF ORGANIC COMPOUNDS

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LABSTRACT

The current knowledge of the gas-phase reactions occurring in the troposphere for organic chemicals is reviewed and evaluated, and areas of uncertainty are discussed.

TABLE OF CONTENTS

		Page
Introd	uction	1
I.	Reactions of Alkyl (R), Alkoxy (RO) and Alkyl Peroxy (RO ₂) Radicals	5
II.	Reactions of Aldehydes, Ketones and $\alpha ext{-Dicarbonyls}$	57
III.	Alkanes	71
IV.	Haloalkanes	81
٧.	Alkenes	84
VI.	Haloalkenes	105
VII.	Alkynes	115
VIII.	Oxygen-Containing Organic Compounds	120
	A. Alcohols	120
	B. Ethers	123
	C. Carboxylic Acids	125
	D. Esters	125
	E. Epoxides	126
	F. Hydroperoxides	127
	G. α,β-Unsaturated Carbonyls	128
	H. 1,4-Unsaturated Carbonyls	132
IX.	Sulfur-Containing Organic Compounds	133
х.	Nitrogen-Containing Organic Compounds	141
XI.	Aromatic Compounds	152
Refere	ences	167

				•	
	*	•			

LIST OF TABLES

Table <u>Number</u>	<u>Title</u>	Page
1	Bimolecular Rate Constants at the High-Pressure Limit k for the Addition Reaction of Alkyl and Substituted-Alkyl Radicals with O ₂ at Around Room Temperature	8
2	Absolute Rate Constant Data for the Reaction of RO ₂ Radicals with NO	15
3	Experimental Rate Constant Ratios $k_a/(k_a+k_b)$ for the Reaction of Secondary Alkyl Peroxy Radicals with NO at ~299 K and 740 Torr Total Pressure of Air	19
4	Absolute Room Temperature Rate Constants for the Gas Phase Reactions of RO ₂ and RCO ₃ Radicals with NO ₂ at the High-Pressure Limit	23
5	Absolute Rate Constant Data for the Gas-Phase Reactions of RO ₂ Radicals with HO ₂	28
6	Kinetic Data for the Gas-Phase Combination Reactions of RO ₂ Radicals	30
7	Absolute Rate Constants for the Reactions of O ₂ with Alkoxy Radicals	40
8	Estimated Arrhenius Parameters and Room Temperature Rate Constants for Decompositions of Alkoxy Radicals	46
9	Estimated Arrhenius Parameters and Room Temperature Rate Constants for 1,4- and 1,5-H Shift Isomerizations of Alkoxy Radicals	52
10	Absolute Rate Constants for Reaction of RO Radicals with NO and NO2	54
11	Absorption Cross Sections, α , and Quantum Yields, ϕ , for the Photolysis of HCHO (from Atkinson et al., 1988a)	59
12	Absorption Cross Section, σ , and Quantum Yields ϕ_a and ϕ_b for CH ₃ CHO photolysis (at 1 atm air) [from Baulch et al., 1984]	61
13	Room Temperature Rate Constants k and Arrhenius Parameters, $k = Ae^{-E/RT}$, for the Reaction of OH Radicals with Carbonyl Compounds. Except as Indicated, Taken from Atkinson (1986)	67
14	Rate Constants at 298 K and Parameters C and D in $k-CT^2e^{-D/T}$ for the Reaction of OH Radicals with Alkanes (from Atkinson 1986a)	73

LIST OF TABLES (continued)

Number	<u>Title</u>	Page
15	Rate Constants for the Reaction of NO_3 Radicals with Alkanes at 296 \pm 2 K (Taken from Atkinson et al., 1988b)	80
16	Room Temperature Rate Constants and Temperature Dependencie $(k = CT^2e^{-D/T})$ for the Gas-Phase Reactions of the OH Radical with Haloalkanes (from Atkinson, 1986a)	
17	Rate Constants k at 298 K and Arrhenius Parameters (k - $Ae^{-E/RT}$) for the Reaction of OH Radicals with Alkanes at the High Pressure Limit (from Atkinson, 1986a)	87
18	Rate Constants k at 298 K and Arrhenius Paramenters ($k = Ae^{-E/RT}$) for the Gas-Phase Reactions of 0_3 with Alkenes, Cycloalkenes and Dialkenes (from Atkinson and Carter, 1984, except as indicated)	96
19	Room Temperature Rate Constants k for the Reaction of NO ₃ Radicals with a Series of Alkenes (from Atkinson et al., 1988b)	102
20	Rate Constants k at Room Temperature and Arrhenius Parameters ($k = Ae^{-E/RT}$) for the Reaction of OH Radicals with Haloalkenes	106
21	Room Temperature Rate Constants k for the Reactions of O ₃ with a Series of Haloalkenes (from Atkinson and Carter, 1984, except as indicated)	111
22	Room Temperature Rate Constants k for the Reactions of NO ₃ Radicals with Haloalkenes (from Atkinson et al., 1988b))114
23	Room Temperature Rate Constants k for the Reaction of OH Radicals with Alkynes at the High-Pressure Limit (from Atkinson, 1986a)	116
24	Room Temperature Rate Constants k and Arrhenius Parameters $(k = Ae^{-E/RT})$ for the Gas-Phase Reactions of the OH Radical with a Series of Alcohols, Ethers, Carboxylic Acids, Esters, Epoxides, Hydroperoxides, α,β -Unsaturated Carbonyls and 1,4-Unsaturated Dicarbonyls	
25	Room Temperature Rate Constants k and Arrhenius Parameters $(k = Ae^{-E/RT})$ for the Gas-Phase Reactions of the OH Radical with Selected Organosulfur Compounds (from Atkinson et al., 1988a, except as indicated)	134

LIST OF TABLES (continued)

Table <u>Number</u>	<u>Title</u>	Page
26	Room Temperature Rate Constants k and Arrhenius Parameters ($k = Ae^{-E/RT}$) for the Gas-Phase Reactions of the NO ₃ Radical with Selected Organosulfur Compounds (from Atkinson et al., 1988a)	139
27	Room Temperature Rate Constants k and Arrhenius Parameters ($k = Ae^{-E/RT}$) for the Gas-Phase Reactions of the OH Radical with Selected Nitrogen-Containing Compounds [from Atkinson (1986a) except as indicated]	143
28	Room Temperature Rate Constants for the Gas-Phase Reactions of the OH Radical with Aromatic Compounds and Rate Constant Ratios $k_a/(k_a + k_b)$	154
29	α-Dicarbonyl Yields from the OH Radical-Initiated Reactions of a Series of Aromatic Hydrocarbons	159
30	Rate Constants for the Gas-Phase Reactions of PAH with OH and NO $_3$ Radicals, N $_2$ O $_5$ and O $_3$ at Room Temperature	160
31	Nitroarene Products Formed from the Gas Phase Reactions of PAH Known to be Present in Ambient Air with OH Radicals (in the Presence of NO_x) and N_2O_5 , Together with their Yields at Room Temperature and Atmospheric Pressure	161

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INTRODUCTION

Large quantities of chemical compounds are emitted into the atmosphere as a result of anthropogenic, biogenic and geogenic processes. These emissions lead to a complex array of chemical and physical transformations resulting in such apparently diverse effects as photochemical air pollution, acid deposition, long-range transport of chemicals, changes in the stratospheric ozone layer and global weather modification. Over the past approximately 15-20 years a vast amount of experimental work, involving laboratory, environmental chamber and ambient atmospheric studies, has been carried out concerning the chemical and physical processes occurring in the atmosphere. Because of the complexity of these processes, the use of computer models is necessary to elucidate and predict the effects of anthropogenic and biogenic emissions on the atmosphere. Such computer models have been used for several years to aid in the formulation of air pollution control strategies, and with the availability of ever more powerful computers, are being used to theoretically investigate the impacts of anthropogenic emissions of reactive organic gases, oxides of nitrogen and sulfur dioxide on the deposition of acidic species in North America and Europe.

Chemical mechanisms, with varying levels of detail, are integral components of these atmospheric computer modeling studies. For tropospheric chemistry applications, such as the modeling of local, urban and regional air pollution impacts and long range transport/acid deposition modeling studies, the detailed chemistry is exceedingly complex, and condensed chemical mechanisms must be utilized. While these detailed and condensed chemical mechanisms are tested against available environmental chamber data during their development, for several model

applications (long range transport, acid deposition modeling and modeling of the clean troposphere) these chamber data are often of limited utility due to the differing organic and oxides of nitrogen concentration regimes and ratios in the environmental chamber experiments compared to those encountered in the modeling scenarios. Thus, the ultimate accuracy of the chemical mechanisms, whether of a detailed or condensed nature, are dependent on the accuracy of the input data, i.e., the rate constants and products of the many hundreds of elementary reactions which occur in the atmosphere.

It is evident that, in parallel with laboratory, ambient atmospheric, and chemical mechanism development and refinement studies, there must also be a continual program to critically evaluate and review the kinetics, mechanisms and reaction products of the chemical reactions relevant to the atmosphere, and to update these evaluations as new experimental and theoretical data become available. For the reactions of importance for chemical modeling of the stratosphere, the National Atmospheric and Space Administration (NASA) established a Panel for Data Evaluation in 1977. To date this panel has published eight, progressively updated and extended, compilations of kinetic and photochemical data for use by modelers in computer simulations of stratospheric chemistry, with the most recent such evaluation appearing in September 1987 (DeMore et al., 1987). In a parallel program, the CODATA Task Group on Chemical Kinetics has critically evaluated the kinetic and photochemical data relevant to middle atmospheric chemistry (10-55 km altitude). The first evaluation appeared in 1980 (Baulch et al., 1980), and two further supplements have been published (Baulch et al., 1982, 1984).

However, these comprehensive evaluations have been concerned primarily with stratospheric modeling (although, obviously, a majority of the reactions covered are also applicable to tropospheric modeling needs), and only organic compounds containing up to three carbon atoms are dealt with (Baulch et al., 1980, 1982, 1984; DeMore et al., 1987). The troposphere, and in particular, the polluted troposphere, contains hundreds of differing organic species of much greater complexity, and there is a need for critical evaluations of the chemical reactions occurring for these more complex organic compounds. Critical review and evaluations of the kinetics and mechanisms of the gas-phase reactions of the hydroxyl (OH) radical and of ozone (O_{Q}) with organic compounds under atmospheric conditions have recently been published (Atkinson and Carter, 1984; Atkinson, 1986a). In addition, and highly relevant to the present evaluation and review, the tropospheric chemistry of eight hydrocarbons (n-butane, 2,3-dimethylbutane, ethene, propene, 1-butene, trans-2-butene, toluene and m-xylene) and their degradation products (Atkinson and Lloyd, 1984) and of the alkanes (Carter and Atkinson, 1985) has been reviewed and evaluated. Despite these extremely useful evaluations, it is evident that there is a need to evaluate the tropospheric chemistry of the various classes of organic compounds emitted into the troposphere from both anthropogenic and biogenic sources, in a manner analogous to the recent evaluation of the atmospheric chemistry of the alkanes of Carter and Atkinson (1985).

This has been carried out in this review, which deals with the chemistry of the major classes of organics of importance in the troposphere. The inorganic reactions which occur in the troposphere are not dealt with here, since these were reviewed and evaluated by Atkinson

and Lloyd (1984) and are the focus of the comprehensive and ongoing NASA and CODATA (now IUPAC) evaluations (Baulch et al., 1980, 1982, 1984; Atkinson et al., 1988a; DeMore et al., 1985, 1987). In this review the previous evaluations noted above are extensively utilized. Thus the recommendations of Atkinson and Carter (1984) and Atkinson (1986a) [updated where necessary] are used for the 03 and OH radical reactions, respectively, and the section dealing with the alkanes is based upon the review of Carter and Atkinson (1985). Since this review and evaluation is an update and extension of the review of Atkinson and Lloyd (1984), extensive use, again updated where necessary, is made of that article whenever possible.

The reactions of organic compounds under tropospheric conditions are dealt with in the following sections:

- (1) reactions of alkyl, alkoxy and alkyl peroxy radicals which are intermediate radical species in the degradation reaction schemes of most organics.
- (2) reactions of aldehydes and ketones, including α -dicarbonyls. These classes of organics are again common "first generation" products formed from the tropospheric degradation reactions of many classes of organics.
 - (3) reactions of alkanes.
 - (4) reactions of haloalkanes.
 - (5) reactions of monoalkenes and di- and trialkenes.
 - (6) reactions of haloalkenes.
 - (7) reactions of alkynes.
- (8) reactions of alcohols, ethers, esters, unsaturated carbonyls, carboxylic acids and hydroperoxides.

- (9) reactions of nitrogen-containing organic compounds.
- (10) reactions of sulfur-containing organic compounds.
- (11) reactions of aromatic hydrocarbons and substituted aromatics, including the polycyclic aromatic hydrocarbons expected to be present in the gas phase.

Only gas-phase processes are discussed; while highly important under many tropospheric conditions, reactions occurring in the particulate and/or aerosol phase (including fog, rain and cloud drops), on surfaces (heterogeneous reactions), and gas- to particle conversion processes are beyond the scope of this review. Clearly, this present review is only one step of a hopefully ongoing critical review process for the atmospheric reactions of organic compounds, and regular updating will be required.

I. REACTIONS OF ALKYL (R), ALKOXY (RO) AND ALKYL PEROXY (RO2) RADICALS

These radicals are common intermediate species formed during the atmospheric degradation reactions of most organic compounds, and their reactions under atmospheric conditions can be treated by class.

Alkyl (R) radicals

Alkyl and substituted alkyl radicals are formed from the initial reactions of several classes of organic compounds with OH radicals, NO_3 radicals and O_3 . For example, alkanes react with OH and NO_3 radicals via H atom abstraction

$$OH + RH \rightarrow H_2O + R$$

$$NO_3 + RH \rightarrow HNO_3 + R$$

and alkenes react with OH and ${\rm NO_3}$ radicals by OH or ${\rm NO_3}$ radical addition to the >C=C< double bond

$$NO_3 + C=C \leftrightarrow C$$

to yield alkyl or β -substituted alkyl radicals. The haloalkanes and haloalkenes react via analogous pathways to form haloalkyl radicals. For the aromatic hydrocarbons, OH radical reaction leads to the initial formation of benzyl and hydroxycyclohexadienyl radicals (or alkylsubstituted homologues), for example, from toluene,

The available kinetic and mechanistic data show that most of these alkyl and substituted alkyl radicals react with $\mathbf{0}_2$ to form an alkyl peroxy radical.

$$R + O_2 \rightarrow RO_2$$

The kinetic data presently available for the O_2 addition pathway are given in Table 1. In addition, Ruiz et al. (1981) and Morgan et al. (1982), have shown that the reaction of O_2 with allyl ($CH_2=CH-CH_2$) radicals proceeds via addition, with the rate constant being in the fall-off region at 413-427 K at total pressures of ≤ 50 torr of argon diluent (Morgan et al., 1982). At approximately 380 K and 50 torr total pressure of argon, the bimolecular rate constant for addition of O_2 to the alkyl radical is 4 x 10^{-13} cm³ molecule⁻¹ s⁻¹ (Morgan et al., 1982).

At elevated temperatures these reactions of alkyl radicals with ${\rm O}_2$ have been assumed to also occur via the H atom abstraction pathway, for example

$$C_2H_5 + O_2 \rightarrow CH_2 = CH_2 + HO_2$$

However, this is now recognized not to be a parallel reaction route, but to occur from the activated ${
m RO}_2$ adduct (Slagle et al., 1984a, 1985; McAdam and Walker, 1987)

$$R + O_2 \neq [RO_2^*] + HO_2 + alkene$$

$$\downarrow M$$

$$RO_2$$

Hence at the high pressure limit at room temperature and below, peroxy radical formation is the sole reaction process. Klais et al. (1979) have also shown that the previously postulated reaction pathway for the reaction of ${\rm CH_3}$ radicals with ${\rm O_2}$

$$CH_3 + O_2 \rightarrow HCHO + OH$$

Table 1. Bimolecular Rate Constants at the High-Pressure Limit k for the Addition Reaction of Alkyl and Substituted-Alkyl Radicals with $\rm O_2$ at Around Room Temperature

Alkyl radical	10 ¹² x k (cm ³ mole- cule ⁻¹ s ⁻¹)	Т (К)	Reference
Methyl	1.8 ± 0.2	298	DeMore et al. (1987)
Ethyl	1.1 ^a 4.4 ± 0.5 4.3 ^a	298 295 295	Plumb and Ryan (1981)
	5.3 ± 0.2 5.0 ± 0.3 4.5 ± 0.2 4.2 ± 0.2 3.7 ± 0.2	298 325 346 376 400	Munk et al. (1986a)
1-Propyl	5.5 ± 0.9	298 ± 3	Ruiz and Bayes (1984)
	5.7	297	Slagle et al. (1985)
2-Propyl	14.1 ± 2.4	298 ± 3	Ruiz and Bayes (1984)
	8.3 ± 0.4	Room temperature	Munk et al. (1986b)
1-Butyl	7.5 ± 1.4	300	Lenhardt et al. (1980)
2-Butyl	16.6 ± 2.2	300	Lenhardt et al. (1980)
2-Methyl-2-propyl	23.4 ± 3.9	300	Lenhardt et al. (1980)
2-Methyl-1-propyl	2.9 ± 0.7	298 ± 3	Wu and Bayes (1986)
2,2-Dimethyl-1-propyl	1.6 ± 0.3	298 ± 3	Wu and Bayes (1986)
Cyclopentyl	17 ± 3	293	Wu and Bayes (1986)
Cyclohexyl	14 ± 2	298 ± 3	Wu and Bayes (1986)
3-Hydroxy-2-butyl	28 ± 18	300	Lenhardt et al. (1980)
сн ₃ co	2.0 ± 0.4	Room temperature	McDade et al. (1982)
			(continued)

Table 1 (continued) - 2

	10 ¹² x k (cm ³ mole-		
Alkyl radical	$cule^{-1} s^{-1})$	T (K)	Reference
с ₆ н ₅ cò	5.7 ± 1.4	Room temperature	McDade et al. (1982)
CF ₃	~8	295	Ryan and Plumb (1982)
	10.0 ± 0.3	300	Cooper et al. (1980)
	9 ± 2	233 - 373	Caralp et al. (1986)
CF ₂ Cl	6.0 ± 1.0 4.7 ^a	298 298	Caralp and Lesclaux (1983)
cc1 ₃	5.1 ± 0.5	300	Cooper et al. (1980)
	~2.5	295	Ryan and Plumb (1984)
Benzyl	0.99 ± 0.77	Room temperature	Ebata et al. (1981)
	1.12 ± 0.11 0.997 ± 0.08 1.03 ± 0.04	295 350 372	Nelson and McDonald (1982)
2-Methylbenzyl	1.2 ± 0.07	Room temperature	Ebata et al. (1981)
4-Methylbenzyl	1.1 ± 0.10	Room temperature	Ebata et al. (1981)

 $^{^{\}rm a}{
m Value}$ at 760 torr total pressure calculated from the fall-off expression.

is of totally negligible importance, having a rate constant of $\langle 3 \times 10^{-16} \rangle$ cm³ molecule⁻¹ s⁻¹ at 368 K.

Hence for the alkyl, haloalkyl, hydroxy-substituted (other than α -hydroxy substituted) alkyl, benzyl and methyl-substituted benzyl and allyl radicals (of general formula $R_1R_2R_3C$ where R_1 , R_2 and R_3 are alkyl, vinyl, halogen or aromatic groups), the reaction with O_2 proceeds via addition to form a peroxy radical, with a room temperature rate constant of $\sim 10^{-12}$ cm³ molecule⁻¹ s⁻¹ at atmospheric pressure. For the smaller alkyl or haloalkyl radicals these reactions are in the fall-off regime between second- and third-order kinetics, but are reasonably close to the high-pressure rate constant at 760 torr of air. Thus, under atmospheric conditions these reactions with O_2 will be the sole loss process of these alkyl type radicals, and other reactions need not be considered.

It is clear that for only a small number of the alkyl-type radicals has the reaction with $\mathbf{0}_2$ been studied, with most proceeding via $\mathbf{0}_2$ addition to form the peroxy radical. However, α -hydroxy radicals, hydroxycyclohexadienyl radicals, and vinyl radicals react by other routes.

The simplest α -hydroxy radical, $\dot{C}H_2OH$, reacts rapidly with O_2 to form the HO_2 radical (Radford, 1980; Wang et al., 1984; Grotheer et al., 1985; Dobé et al., 1985)

with a room temperature rate constant of (in units of 10^{-12} cm³ molecule⁻¹ s⁻¹) 2^{+2}_{-1} (Radford, 1980); 1.4 ± 0.4 (Wang et al., 1984), 9.5 ± 2.5 (Grotheer et al., 1985), 10.6 ± 2.5 (Dobé et al., 1985). The most recent data are to be preferred (Demore et al., 1987), with a rate constant of

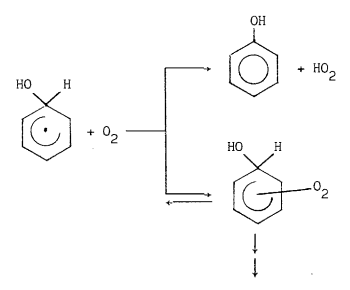
9.6 x 10^{-12} cm³ molecule⁻¹ s⁻¹ at 298 K being recommended. With a rate constant of this magnitude, reaction with 0_2 will be the sole loss process of CH₂OH radicals in the atmosphere. Product studies have shown that the higher (C_2-C_4) α -hydroxy radicals also react via H-atom abstraction to yield the corresponding carbonyls (Carter et al., 1979a; Ohta et al., 1982)

$$R_1 \rightarrow COH + O_2 \rightarrow RCOR_1 + HO_2$$

As discussed by Atkinson and Lloyd (1984) and Atkinson (1986a), the expected radical resulting from O_2 addition to $\dot{C}H_2OH$, $\dot{O}OCH_2OH$, is also presumably formed from the reaction of HO_2 radicals with HCHO

This suggests that the exothermicity of the O_2 + CH_2OH reaction leads to rapid decomposition of the energy-rich $OOCH_2OH$ radical (and other $RR_1C(OH)OO$ radicals).

Hydroxycyclohexadienyl and methyl-substituted hydroxycyclohexadrenyl radicals are formed from OH radical addition to the aromatic hydrocarbons. As discussed in Section (XI) below, the subsequent reactions of these hydroxycyclohexadienyl radicals lead to formation of phenols and to aromatic ring cleavage. These observed products suggest that reaction with O_2 occurs (see, for example, Atkinson and Lloyd, 1984)



ring cleavage products.

However, Zellner et al. (1985) have directly observed that while the hydroxycyclohexadienyl radical reacts rapidly with NO and NO_2 at 298 K,

$$C_6H_6OH + NO \rightarrow products$$

 $k = (1.0 \pm 0.5) \times 10^{-12} \text{ cm}^3 \text{ molecule}^{-1} \text{ s}^{-1}$

$$C_6H_6OH + NO_2 \rightarrow products$$

 $k = (8.5 \pm 2.1) \times 10^{-12} \text{ cm}^3 \text{ molecule}^{-1} \text{ s}^{-1}$

no observable reaction with 0_2 was found (with a rate constant of $\le 10^{-13}$ cm³ molecule⁻¹ s⁻¹ being suggested by their data). However, this upper limit to the 0_2 reaction rate constant does not preclude the 0_2 reaction being dominant under atmospheric conditions. More recent evidence suggesting that these 0_2 reactions are not important under atmospheric conditions arises from the hydroxyaromatic and nitroarene data of Atkinson et al. (1987a) for naphthalene and biphenyl (see Section XI).

Slagle et al. (1984b) have investigated the kinetics of the reaction of the vinyl radical with 0_2 , and observed that the rate constant is pressure independent, with

$$k = 6.6 \times 10^{-12} e^{126/T} cm^3 molecule^{-1} s^{-1}$$

over the temperature range 297-602 K. The magnitude of this rate constant, and the almost zero temperature dependence, strongly suggest the initial formation of a ${\rm C_2H_3O_2}$ adduct. However, the lack of a pressure dependence and the observation of HCHO and HCO radicals as products of this reaction show that the reaction proceeds via

$$^{\circ}C_{2}H_{3} + O_{2} \rightarrow [H_{2}C = CHOO] \rightarrow HCHO + HCO$$

Presumably the homologues of the vinyl radical react analogously.

Alkyl peroxy (RO₂) radicals

These radicals are formed from the addition of 0_2 to the alkyl radicals, as discussed above:

$$R + O_2 \rightarrow RO_2$$

Under tropospheric conditions, RO_2 radicals react with NO, via two pathways,

$$RO_2 + NO \xrightarrow{M} RONO_2$$
 (a)

with HO_2 radicals,

$$RO_2 + HO_2 \rightarrow ROOH + O_2$$
(and possibly other products)

and RO_2 radicals (either a self-reaction or a reaction with other alkyl peroxy radicals)

$$RO_2 + RO_2 \rightarrow \text{products}.$$

The reaction pathways which occur depend on the NO to HO_2 and/or RO_2 radical concentration ratios; and in the troposphere reaction with NO is expected to dominate for NO mixing ratios $\gtrsim 30$ ppt (Logan et al., 1981; Logan, 1983). The reaction of RO_2 radicals with NO_2

$$RO_2 + NO_2 \rightarrow RO_2NO_2$$

are generally unimportant under tropospheric conditions due to the rapid thermal decomposition of the alkyl peroxynitrates back to reactants.

These reactions are discussed in the following sections.

Reaction with NO

The kinetic data obtained using absolute measurement techniques are given in Table 2. The studies of Cox and Tyndall (1979, 1980), Sander and Watson (1980), Ravishankara et al. (1981), Simonaitis and Heicklen (1981) and Plumb et al. (1981) for the CH₃O₂ reaction appear to be free of possible experimental problems and are in good agreement. Based largely upon these studies NASA (DeMore et al., 1987) and CODATA (Baulch et al.,

Table 2. Absolute Rate Constant Data for the Reaction of RO_2 Radicals with NO

	10 ¹² x k (cm ³ mole-	, ₍₁₀	
R	$cule^{-1} s^{-1})$	T (K)	Reference
CH ₃ 0 ₂	>1	а	Anastasi et al. (1978)
	8.0 ± 2.0	295 ± 2	Plumb et al. (1979)
	2.92 ± 0.12	а	Adachi and Basco (1979a)
	6.5 ± 2.0	298	Cox and Tyndall (1979,1980)
	7.1 ± 1.4	298	Sander and Watson (1980)
	8.4 ± 1.5 8.6 ± 1.1 9.0 ± 1.1 7.8 ± 1.2 7.8 ± 1.4	240 250 270 298 339	Ravishankara et al. (1981)
	13.5 ± 1.4 17.0 ± 2.2 7.7 ± 0.9 6.3 ± 0.5	218 218 296 365	Simonaitis and Heicklen (1981)
	8.6 ± 2.0	295	Plumb et al. (1981)
	7 ± 2	298	Zellner et al. (1986)
сн ₃ сн ₂ 0 ₂	2.66 ± 0.17	а	Adachi and Basco (1979b)
	8.9 ± 3.0	295	Plumb et al. (1982)
(CH ₃) ₂ CHO ₂	3.5 ± 0.3	a	Adachi and Basco (1982a)
(CH ₃) ₃ CO ₂	>1	a	Anastasi et al. (1978)
CF ₃ O ₂	14.5 ± 2.0	298 ^b	Dognon et al. (1985)
	17.8 ± 3.6	295	Plumb and Ryan (1982)
CF ₂ ClO ₂	16 ± 3	298 ^b	Dognon et al. (1985)
CFC1 ₂ 0 ₂	16 ± 2	a	Lesclaux and Caralp (1984)
	14.5 ± 2.0	298 ^b	Dognon et al. (1985)
			(continued)

Table 2 (continued) - 2

R	10^{12} x k $(\text{cm}^3 \text{ mole-}$ $\text{cule}^{-1} \text{ s}^{-1})$	T (K)	Reference	
cc1 ₃ 0 ₂	18.6 ± 2.8	295	Ryan and Plumb (1984)	
	17.0 ± 2.0	298 ^b	Dognon et al. (1985)	

aRoom temperature, not reported. bStudied over the temperature range ~230-430 K, individual rate constants other than at 298 K not reported.

1984) recommended that for the reaction of CH_3O_2 radicals with NO

$$k(CH_3O_2 + NO) = 4.2 \times 10^{-12} e^{(180 \pm 180)/T} cm^3 molecule^{-1} s^{-1},$$

= 7.6 x 10⁻¹² cm³ molecule⁻¹ s⁻¹ at 298 K,

This rate constant expression was also recommended by Atkinson and Lloyd (1984). The rate constant determined by Plumb et al. (1982) for ${\rm C_2H_5O_2}$ is, within the cited experimental errors, identical to that for the ${\rm CH_3O_2}$ radical.

Hence, consistent with the treatment of Atkinson and Lloyd (1984), the rate constant for the reaction of all alkyl peroxy (RO_2) and acyl peroxy (RCO_3) radicals with NO is recommended to be

$$k(RO_2 + NO) = k(RCO_3 + NO)$$

= $4.2 \times 10^{-12} e^{180/T} cm^3 molecule^{-1} s^{-1}$
= $7.6 \times 10^{-12} cm^3 molecule^{-1} s^{-1} at 298 K$

The reaction of CH_3O_2 with NO has been shown to proceed primarily via the reaction

$$CH_3O_2 + NO + CH_3O + NO_2$$

(Pate et al., 1974; Ravishankara et al., 1981; DeMore et al., 1987; Zellner et al., 1986), and Plumb et al. (1982) have shown that the reaction of $\rm C_2H_5O_2$ radicals with NO yields NO₂ with a yield of $\rm \geq 0.80$.

However, for the larger alkyl peroxy radicals, Darnall et al. (1976), Atkinson et al. (1982, 1983a, 1984a, 1987b) and Takagi et al. (1981) have shown that the reaction pathway

$$RO_2 + NO \rightarrow [ROONO] \rightarrow RONO_2$$

becomes important. The product data of Atkinson et al. (1982, 1983a, 1984a, 1987b) show that for the secondary alkyl peroxy radicals the rate constant ratio $k_a/(k_a + k_b)$, where k_a and k_b are the rate constants for the reaction pathways (a) and (b), respectively, at room temperature and atmospheric pressure

$$RO_{2} + NO \longrightarrow RO + NO_{2}$$

$$RO_{2} + NO \longrightarrow RO + NO_{3}$$

$$(a)$$

increases monotonically with the carbon number of the RO_2 radical (Table 3). Furthermore, for a given alkyl peroxy radical the rate constant ratio $k_a/(k_a+k_b)$ is pressure—and temperature—dependent, increasing with increasing pressure and with decreasing temperature (Atkinson et al., 1983a, 1987b). The pressure and temperature—dependent rate constant ratios k_a/k_b for secondary alkyl peroxy radicals are fit by the fall-off expression (Carter and Atkinson, 1988) [note that earlier evaluations of Atkinson et al. (1983a), Carter and Atkinson (1985) and Atkinson (1986) utilized the ratio $k_a/(k_a+k_b)$ rather than k_a/k_b (see Carter and Atkinson, 1988)].

Table 3. Experimental Rate Constant Ratios $\rm k_a/(k_a+k_b)$ for the Reaction of Secondary Alkyl Peroxy Radicals with NO at ~299 K and 740 Torr Total Pressure of Air

Carbon Number	k _a /(k _a + k _b) ^a	Peroxy Radical	
3	0.042 ± 0.003	2-Propyl	
4	0.090 ± 0.008	2-Butyl	
5	0.129 ± 0.016, 0.134 ± 0.002	2-Pentyl	
	0.131 ± 0.016 , 0.146 ± 0.009	3-Pentyl	
	0.141 ± 0.003 , 0.150 ± 0.004	2-Methyl-3-butyl	
6	0.209 ± 0.032	2-Hexyl	
	0.230 ± 0.031	3-Hexyl	
	0.160 ± 0.015	Cyclohexyl	
	0.190 ± 0.018	2-Methyl-3-pentyl	
		+ 2-Methyl-4-pentyl	
·	0.162 ± 0.009 , 0.178 ± 0.017	3-Methyl-2-pentyl	
7	0.291 ± 0.022, 0.301 ± 0.049	2-Heptyl	
	0.323 ± 0.048, 0.325 ± 0.014	3∸Heptyl	
	0.285 ± 0.015 , 0.301 ± 0.045	4-Heptyl	
8	0.323 ± 0.024	2.0-41	
O		2-0ety1	
	0.348 ± 0.032	3-0ety1	
	0.329 ± 0.032	4-Octyl	

^aAt 298-300 K and 735-740 torr total pressure (from Atkinson et al. 1982, 1983a, 1984a, 1987b). The indicated error limits are two least-squares standard deviations, but do not take into account errors due to GC-FID calibrations of the alkanes and alkyl nitrates, nor of the estimation procedure used to derive the rate constant ratios from the observed alkyl nitrate yields.

$$\frac{k_{a}}{k_{b}} = \left(\frac{Y_{o}^{300}[M](T/300)^{-m}o}{Y_{o}^{300}[M](T/300)^{-m}o}\right) F^{2}$$

$$\left(1\right)$$

where

$$z = \left\{ 1 + \left[log \left(\frac{Y_o^{300}[M](T/300)^{-m}o}{Y_\infty^{300}(T/300)^{-m}\omega} \right) \right]^2 \right\}^{-1}$$

and $Y_0^{300} = \alpha e^{\beta n}$, where n is the number of carbon atoms in the alkyl peroxy radical and α and β are constants. The most recent evaluation of the experimental data, (Carter and Atkinson, 1988) leads to

$$Y_{\infty}^{300} = 0.826$$
 $\alpha = 1.94 \times 10^{-22} \text{ cm}^3 \text{ molecule}^{-1}$
 $\beta = 0.97$
 $m_0 = 0$
 $m_{\infty} = 8.1$

F = 0.411

and

Although the rate constant ratios k_a/k_b at room temperature and atmospheric pressure for secondary RO_2 radicals depend primarily on the number of carbon atoms in the RO_2 molecule, the corresponding rate constant ratios for primary and tertiary RO_2 radicals are significantly

lower, by a factor of ~2.5 for primary and a factor of ~3.3 tertiary alkyl peroxy radicals (Atkinson et al., 1987b; Carter and Atkinson, 1988).

Accordingly,

$$(k_a/k_b)$$
 primary $\approx 0.40 (k_a/k_b)$ secondary

and

$$(k_a/k_b)$$
 tertiary = 0.3 (k_a/k_b) secondary

It should be noted that the use of the above equations to calculate rate constant ratios k_a/k_b is solely applicable to alkyl peroxy radicals. Thus, although no definitive data exist, computer modeling data suggest (Carter and Atkinson, 1985) that the rate constant ratios for δ -hydroxyalkyl peroxy radicals (for example RCHOHCH₂CH₂CH(00)R⁻) are much lower than for the corresponding alkyl peroxy radicals.

These alkyl nitrate forming reactions are postulated to occur via the mechanism (Atkinson et al., 1983a)

For the haloalkyl peroxy radicals CX_3O_2 , where X=F or Cl, the room temperature reaction rate constants are all ~(1.5-1.9) x 10^{-12} cm³ molecule⁻¹ s⁻¹ (Table 2), a factor of ~2 higher than those for the alkyl peroxy radicals. Dognon et al. (1985) have determined rate constants for the reactions of CF_3O_2 , CF_2ClO_2 , CFCl_2O_2 and CCl_3O_2 with NO over the temperature range ~230-430 K. Negative temperature dependences were observed, with the values of n in the expression k = k₂₉₈ (T/298)⁻ⁿ being 1.2 ± 0.2, 1.5 ± 0.4, 1.3 ± 0.2 and 1.0 ± 0.2 for CF_3O_2 , CF_2ClO_2 , CFCl_2O_2 and CCl_3O_2 , respectively. These negative temperature dependences are equivalent, over this temperature range, to an Arrhenius activation energy of -720 cal mole⁻¹. Use of this Arrhenius activation energy leads to

$$k(CX_3O_2 + NO; X = F, C1) = 5.0 \times 10^{-12} e^{360/T} cm^3 molecule^{-1} s^{-1}$$

$$= 1.67 \times 10^{-11} cm^3 molecule^{-1} s^{-1} at 298 K$$

which is recommended in the absence of further data. The available data (Lesclaux and Caralp, 1984; Dognon et al., 1985) suggest that these reactions of the haloalkylperoxy radicals with NO proceed via

$$CX_3O_2 + NO \rightarrow CX_3O + NO_2$$

Reaction with NO2

The kinetic data obtained using absolute experimental methods are given in Table 4. The most recent studies of Sander and Watson (1980) and Ravishankara et al. (1980) for ${\rm CH_3O_2}$ show that this reaction is in the fall-off regime between second- and third-order kinetics below atmospheric

Table 4. Absolute Room Temperature Rate Constants for the Gas Phase Reactions of RO_2 and RCO_3 Radicals with NO_2 at the High-Pressure Limit

RO ₂	10^{12} x k (cm ³ molecule ⁻¹ s ⁻¹)	Reference
CH ₃ 0 ₂	1.6 ± 0.3 ^a (540 torr N ₂)	Cox and Tyndall (1979, 1980)
	1.53 ± 0.07 (53-580 torr 0 ₂ + Ar)	Adachi and Basco (1980)
	8.0 ± 1.0	Sander and Watson (1980)
	7	Ravishankara et al. (1980)
с ₂ н ₅ о ₂	1.25 ± 0.07	Adachi and Basco (1979c)
(CH ₃) ₂ CHO ₂	5.65 ± 0.17	Adachi and Basco (1982a)
(CH ₃) ₃ CO ₂	≥0.5	Anastasi et al. (1978)
CFCl ₂ O ₂	6.0 ± 1.0	Lesclaux and Caralp (1984)
CH ₃ C(0)0 ₂	6.0 ± 2.0 (714 torr 0 ₂)	Addison et al. (1980)
	6.1	Basco and Parmar (1987)

^aAt 275 K.

thermal decomposition data of Reimer and Zabel (1986). The data of Adachi and Basco (1980) appear to have been in error due to interferences by the ${\rm CH_3O_2NO_2}$ product with the spectroscopic technique used, and this was probably also the case for their studies of the ${\rm C_2H_5O_2}$ and ${\rm (CH_3)_2CHO_2}$ radical reactions.

For $\mathrm{CH_{3}O_{2}}$, Ravishankara et al. (1980) fit the data for $\mathrm{M=N_{2}}$ from their study and that of Sander and Watson (1980), which are in excellent agreement, to the fall-off expression

$$k = \left(\frac{k_{o}(T)[M]}{1 + k_{o}(T)[M]}\right) F \left\{1 + [\log (k_{o}(T)[M]/k_{\omega}(T)]^{2}\right\}^{-1}$$
 (I) with

E - 0

$$k_0(T) = 2.2 \times 10^{-30} (T/298)^{-2.5} \text{ cm}^6 \text{ molecule}^{-2} \text{ s}^{-1}$$

and

$$k_m(T) = 7 \times 10^{-12} (T/298)^{-3.5} \text{ cm}^3 \text{ molecule}^{-1} \text{ s}^{-1}$$

Hence $k = 3.7 \times 10^{-12} \text{ cm}^3 \text{ molecule}^{-1} \text{ s}^{-1} \text{ at 298 K and 760 torr total}$ pressure.

This expression, which was recommended by Atkinson and Lloyd (1984), is slightly different to the NASA (DeMore et al., 1987) and CODATA (Baulch et al., 1982) evaluations, but gives a better fit to the experimental data. For the $\geq C_2$ alkyl peroxy radicals, apart from the lower limit to the rate constants for (CH₃)₃CO₂ derived by Anastasi et al. (1978), only the suspect data of Adachi and Basco (1979c, 1982a) are available. Atkinson and Lloyd (1984) recommended that the limiting high-pressure rate

constants for the $\geq C_2$ alkyl peroxy radicals are identical to that for ${
m CH_3O_2},$

$$k_{\infty}(RO_2 + NO_2) = 7 \times 10^{-12} (T/298)^{-3.5} \text{ cm}^3 \text{ molecule}^{-1} \text{ s}^{-1}$$

The pressures at which these reactions will exhibit kinetic fall-off behavior from the second- to third-order regime will decrease as the size of the RO_2 radical increases, and it is expected that at room temperature and 760 torr total pressure these $\ge C_2$ alkyl peroxy reactions will be close to the limiting high-pressure region.

For the acyl peroxy (RCO $_3$) radicals, rate constants are available only for the CH $_3$ C(0)00 radical (Addison et al., 1980; Basco and Parmar, 1987). Both studies, which are in good agreement, show that the rate constant is pressure dependent [in agreement with the data of Reimer and Zabel (1986) on the thermal decomposition of CH $_3$ C(0)00NO $_2$]. Basco and Parmar (1987) have fitted their data (obtained at room temperature) to the Troe fall-off equation, and obtain:

$$F = 0.19$$
 $k_0 = 5.07 \times 10^{-29} \text{ cm}^6 \text{ molecule}^{-2} \text{ s}^{-1}$ and

$$k_{\infty} = 6.09 \times 10^{-12} \text{ cm}^3 \text{ molecule}^{-1} \text{ s}^{-1}$$

This high-pressure limit is similar to that for the ${
m RO_2}$ + ${
m NO_2}$ reactions, and it is recommended that for all ${
m RCO_3}$ and ${
m RO_2}$ radical reactions have the same limiting high pressure rate constant at 298 K of

$$k(RO_2 + NO_2; RCO_3 + NO_2) = 7 \times 10^{-12} cm^3 molecule^{-1} s^{-1}$$

For $CFCl_2O_2$, Lesclaux and Caralp (1984) and Lesclaux et al. (1986) fit their room temperature data to equation (I) with

$$F = 0.6$$

$$k_0 = (3.5 \pm 0.5) \times 10^{-29} \text{ cm}^6 \text{ molecule}^{-2} \text{ s}^{-1}$$
and
$$k_m = (6.0 \pm 1.0) \times 10^{-12} \text{ cm}^3 \text{ molecule}^{-1} \text{ s}^{-1}$$

This reaction is hence reasonably close to the high pressure limit at 298 K and 760 torr total pressure (k = 5.4 x 10^{-12} cm³ molecule⁻¹ s⁻¹). In a recent temperature dependence study, Lesclaux et al. (1986) fit their data for CFCl₂O₂ with k_o = (3.5 ± 0.5) x 10^{-29} (T/298)^{-(4.1 ± 0.3)} cm⁶ molecule⁻² s⁻¹ and k_∞ = (5'.9 ± 1.0) x 10^{-12} (T/298)^{-(3.6 ± 0.5)} cm³ molecule⁻¹ s⁻¹. In the absence of further experimental data this expression should be used for the other CX₃O₂ (X = F or Cl) radicals.

These absolute rate data are supported by the relative rate data of Simonaitis et al. (1979) and Simonaitis and Heicklen (1979) for $CFCl_2O_2$ and CCl_3O_2 radicals, for which the rate constant ratios $k(CX_3O_2 + NO_2)/k(CX_3O_2 + NO) = 0.58 \pm 0.10$ ($CFCl_2O_2$) and 0.68 (CCl_3O_2) were obtained at 1 atmosphere pressure of N_2 , independent of temperature over the range 268-298 (CCl_3O_2) and 286-305 K ($CFCl_2O_2$). Thus the limiting high pressure rate constants for the reactions of CX_3O_2 radicals with NO_2 are similar to those for RO_2 and RCO_3 radicals.

These reactions of alkyl peroxy and haloalkyl peroxy radicals with NO2 all proceed via combination to yield the corresponding peroxynitrates

$$RO_2 + NO_2 \rightarrow ROONO_2$$

(Niki et al., 1978a, 1980; Edney et al., 1979; Spence et al., 1978; Hendry and Kenley, 1979; Gay et al., 1976; Simonaitis and Heicklen, 1979; Simonaitis et al., 1979).

Reaction with HO2 Radicals

Relatively few data exist for the reactions of ${\rm HO_2}$ radicals with ${\rm RO_2}$ radicals. Cox and Tyndall (1979, 1980) used molecular modulation spectroscopy to study the reaction of ${\rm HO_2}$ radicals with ${\rm CH_3O_2}$ over the temperature range 274-337 K. Assuming, in agreement with Sander and Watson (1981), that the ${\rm CH_3O_2}$ radical absorption cross-section is independent of temperature, with $\sigma({\rm CH_3O_2}) = 3.9 \times 10^{-18} \ {\rm cm^2}$ at 250 nm, the rate constants given in Table 5 were obtained. A product was observed which adsorbed strongly in the wavelength region 210-280 nm, and attributed to ${\rm CH_3OOH}$. At ${\rm CM_3OOH}$ at spectrum of this product was reasonably similar to that reported by Molina and Arguello (1979), supporting the conclusion that the reaction proceeds via

$$CH_3O_2 + HO_2 + CH_3OOH + O_2$$

More recently, McAdam et al. (1987) and Kurylo et al. (1987) have determined the absorption cross-sections of ${\rm CH_{3}O_{2}}$ (and ${\rm HO_{2}}$) and also measured the rate constant for the reaction of ${\rm CH_{3}O_{2}}$ with ${\rm HO_{2}}$ at 298 K (Table 5).

Table 5. Absolute Rate Constant Data for the Gas-Phase Reactions of RO_2 Radicals with HO_2

			
RO ₂	10^{12} x k $(\text{cm}^3 \text{ molecule}^{-1} \text{ s}^{-1})$	T (K)	Reference
сн ₃ 0 ₂	8.5 ± 1.2 6.5 ± 1.0 3.5 ± 0.5	274 298 338	Cox and Tyndall (1980)
	6.4 ± 1.0	298	McAdam et al. (1987)
	2.9 ± 0.4	298	Kurylo et al. (1987)
С ₂ H ₅ O ₂	6.3 ± 0.9	295	Cattrell et al. (1986)
	7.3 ± 1.0 6.0 ± 0.5 5.3 ± 1.0 3.4 ± 1.0 3.1 ± 0.5	248 273 298 340 380	Dagaut et al. (1988a)

For the reaction of ${\rm C_2H_5O_2}$ with ${\rm HO_2}$, kinetic data have recently been reported by Cantrell et al. (1986) and Dagaut et al. (1988a). The rate constants derived in these studies depend on the absorption cross-sections measured or assumed for the ${\rm HO_2}$ and ${\rm RO_2}$ radicals. Furthermore, the ultraviolet absorption bands of these species overlap, resulting in the potential for significant overall uncertainties to be associated with the reported measurements of these radical-radical reaction rate constants.

The data given in Table 5 show that while the reported 298 K rate constants for the reaction of HO_2 radicals with $\mathrm{CH}_3\mathrm{O}_2$ radicals vary by a factor of 2, those for the reaction of HO_2 radicals with $\mathrm{C}_2\mathrm{H}_5\mathrm{O}_2$ radicals are in good agreement. Rather than recommend a specific study, a rate constant at 298 K for the reactions of HO_2 radicals with RO_2 radicals of

$$k(HO_2 + RO_2) = 5 \times 10^{-12} \text{ cm}^3 \text{ molecule}^{-1} \text{ s}^{-1}$$

is indicated, with a likely overall uncertainty of \pm a factor of 2. The temperature dependent studies for both of the reactions studied to date yield negative temperature dependencies equivalent to E/R values of -1300 K (Cox and Tyndall, 1980) and -650 K (Dagaut et al., 1988a). A value of E/R = -1000 K is chosen to yield

$$k(HO_2 + RO_2) = 1.75 \times 10^{-13} e^{1000/T} cm^3 molecule^{-1} s^{-1}$$

Reaction with RO2 Radicals

Numerous studies of the self reactions of RO₂ radicals have been carried out, and the data obtained are listed in Table 6. These data are discussed below:

Table 6. Kinetic Data for the Gas-Phase Combination Reactions of RO_2 Radicals

RO ₂	$10^{13} \text{ x k}_{\text{obs}}$ (cm ³ molecule ⁻¹ s ⁻¹)	т (К)	Reference
СH ₃ 0 ₂	4.0	298	DeMore et al. (1987) ^a
С ₂ Н ₅ О ₂	1.10 ± 0.09 1.0 ± 0.1 ^b	Room temperature	Adachi et al. (1979)
	0.948 ± 0.130 1.10 ± 0.15 1.33 ± 0.19 1.51 ± 0.21 1.68 ± 0.23	303 333 373 423 457	Anastasi et al. (1983)
	0.52	298	Munk et al. (1986a)
	1.30 ± 0.16 (0.84 ^b)	268-347.5	Cattell et al. (1986)
	0.59 ^c	298	Dagaut et al. (1988a)
сн ₃ сн ₂ сн ₂ о ₂	3.84 ± 0.33	Room temperature	Adachi and Basco (1982)
(сн ₃) ₂ сно ₂	0.0135 ± 0.0008 0.0181 ± 0.0020 0.0269 ± 0.0020 0.0418 ± 0.0050 0.0591 ± 0.0033	300 313 333 353 373	Kirsch et al. (1978)
	0.0203 ± 0.0058	Room temperature	Adachi and Basco (1982)
	0.53 ± 0.05	Room temperature	Munk et al. (1986b)
(CH ₃) ₃ CO ₂	0.000256 0.00088	298 325	Anastasi et al. (1978)
	0.000256	Room temperature	Parkes (1975)
	0.000188 ^c	298	Kirsch et al. (1978)
CH ₃ C(0)0 ₂	25 ± 10	302 ± 1	Addison et al. (1980)
	80 ± 15	Room temperature	Basco and Parmar (1985

aEvaluation.
bEstimated rate constant for the elementary reaction (see text).
cCalculated from cited Arrhenius expression.

$CH_3O_2 + CH_3O_2 \rightarrow products$

The majority of kinetic and product studies concerning the self-reactions of RO_2 radicals have concerned this reaction. However, since the rate constants have been derived from measurements of k/σ and σ , where σ is the absorption cross section, and most studies were carried out at different wavelengths, comparison of the reported data is somewhat difficult. Absorption spectra and cross-sections have been reported from several studies. The most recent NASA evaluation (DeMore et al., 1987) used the absorption cross-sections determined by Hochanadel et al. (1977) to reevaluate the room temperature literature rate constants, yielding

$$k_{obs}(CH_3O_2 + CH_3O_2) = 4.0 \times 10^{-13} \text{ cm}^3 \text{ molecule}^{-1} \text{ s}^{-1} \text{ at 298 K,}$$

with an estimated overall uncertainty of \pm a factor of 1.5, where k_{obs} is the experimentally measured rate constant which may include contributions from secondary reactions. Combined with the temperature dependence of Sander and Watson (1981), this leads to (DeMore et al., 1987)

$$k_{obs}(CH_3O_2 + CH_3O_2) = 1.9 \times 10^{-13} e^{220/T} cm^3 molecule^{-1} s^{-1}$$

Kan and Calvert (1979) and Kurylo et al. (1987) have shown that, in contrast to the combination reaction of ${\rm HO_2}$ radicals, ${\rm H_2O}$ vapor has no effect on the measured rate constant for combination of ${\rm CH_3O_2}$ radicals at room temperature.

However, secondary reactions of the products of this reaction will result in these measured values being upper limits to the true elementary rate constant. The reaction proceeds via the pathways:

$$CH_{3}O_{2} + CH_{3}O_{2} \rightarrow 2 CH_{3}O + O_{2}$$
 (a)

$$\rightarrow$$
 HCHO + CH₃OH + O₂ (b)

$$+ CH_3O_2CH_3 + O_2$$
 (c)

Quantitative data for these reaction pathways have been obtained by Parkes (1975), using molecular modulation spectroscopy, and more recently by Kan et al. (1980) and Niki et al. (1981a) using FT-IR absorption spectroscopy. These data are in good agreement and Atkinson and Lloyd (1984) recommend

$$k_a/k_{overall} = 0.35$$

$$k_b/k_{overall} = 0.57$$

and

$$k_c/k_{overall} = 0.08$$

with the ratios $k_a/k_{overall}$ and $k_c/k_{overall}$ being uncertain by $\pm 30\%$. Hence, secondary reactions of CH₃O radicals, for example

$$CH_{3}O + CH_{3}O_{2} \rightarrow CH_{3}OOH + HCHO$$

$$CH_3O + O_2 \rightarrow HCHO + HO_2$$

$$CH_3O_2 + HO_2 \rightarrow products$$

lead to the observed rate constant being an upper limit to the "true" rate constant. Kan et al. (1980) have assessed the influence of secondary reactions in this system and estimate that the value of $k_{\rm overall}$ is some 12% lower than the observed rate constant. Thus, it is recommended that

$$k(CH_3O_2 + CH_3O_2) = 3.5 \times 10^{-13} \text{ cm}^3 \text{ molecule}^{-1} \text{ s}^{-1}$$

at 298 K, and

$$k(CH_3O_2 + CH_3O_2) = 1.7 \times 10^{-13} e^{220/T} cm^3 molecule^{-1} s^{-1}$$

$$C_2H_5O_2 + C_2H_5O_2 \rightarrow Products$$

Room temperature rate constants have been determined for this reaction by Adachi et al. (1979), Anastasi et al. (1983), Munk et al. (1986a), Cattell et al. (1986) and Dagaut et al. (1988a) and, as seen from Table 6, the measured rate constants range over a factor of ~2. The reported temperature dependencies are zero (Cattell et al., 1986) or slightly positive (Anastasi et al., 1983; Dagaut et al., 1988a). This reaction can proceed via the channels

$$C_2^{H_5}O_2 + C_2^{H_5}O_2 \rightarrow 2 C_2^{H_5}O + O_2$$
 (a)

$$+ C_2 H_5 O \dot{H} + C H_3 C H O + O_2$$
 (b)

$$+ C_2 H_5 OOC_2 H_5 + O_2$$
 (c)

Niki et al. (1982a) obtained the rate constant ratios $k_a/k_b=1.3\pm0.16$ and $k_c/k_b\le0.22$, which leads to $k_a/k_{overall}=0.52$ to 0.56, $k_b/k_{overall}=0.40$ to 0.44, and $k_c/k_{overall}\le0.09$. Anastasi et al. (1983) determined that $k_a/k_b=1.75\pm0.05$ at 302 K, 2.12 \pm 0.10 at 333 K, and 2.45 \pm 0.15 at 373 K, with channel (c) contributing 5% of the overall reaction.

Using the recent kinetic data of Cattell et al. (1986) with k_a/k_b = 1.5 and k_c/k_a ~ 0 yields the elementary rate constants, corrected for secondary reactions, of

$$k_a = 5 \times 10^{-14} \text{ cm}^3 \text{ molecule}^{-1} \text{ s}^{-1}$$

and

$$k_b = 3.5 \times 10^{-14} \text{ cm}^3 \text{ molecule}^{-1} \text{ s}^{-1}$$

both independent of temperature over the small range of ~270-350 K.

$$\underline{\text{CH}_3\text{CH}_2\text{CH}_2\text{O}_2} + \underline{\text{CH}_3\text{CH}_2\text{CH}_2\text{O}_2} \rightarrow \underline{\text{Products}}$$

The sole reported data concerning this reaction are those of Adachi and Basco (1982b), who obtained a value of

$$k_{obs} = (3.8 \pm 0.3) \times 10^{-13} \text{ cm}^3 \text{ molecule}^{-1} \text{ s}^{-1}$$

at room temperature. No determination of the products formed were made, and this rate constant is an upper limit to the "true" rate constant for this reaction.

$\frac{(CH_3)_2CHO_2. + (CH_3)_2CHO_2. \rightarrow Products}{(CH_3)_2CHO_2. \rightarrow Products}$

This reaction has been investigated by Kirsch et al. (1978, 1979), Cowley et al. (1982), Adachi and Basco (1982b), and Munk et al. (1986b), with the only temperature dependence study being that of Kirsch et al. (1978) over the range 300-373 K using molecular modulation spectroscopy. The rate constant reported by Munk et al. (1986b) is over an order of magnitude higher than the other data given in Table 6. One wonders if it is possible that they were not studying only the reactions of the isopropylperoxy radical.

These observed rate constants are again upper limits to the "true" rate constant because of the influence of secondary reactions of the isopropoxy radicals formed in reaction (b)

$$2(CH_3)_2CHO_2 \rightarrow (CH_3)_2CHOH + (CH_3)_2CO + O_2$$
 (a)

$$+ 2(CH_3)_2CHO + O_2$$
 (b)

and Kirsch et al. (1979) and Cowley et al. (1982) obtained, from a product analysis study, the rate constant ratios

$$k_b/k_a = 1.39 \pm 0.04$$
 at 302 K

$$= 1.84 \pm 0.04 t 333 K$$

$$= 2.80 \pm 0.08$$
 at 373 K

From these data and the observed rate constants of Kirsch et al. (1978), the Arrhenius expressions for reaction pathways (a) and (b) were derived (Cowley et al., 1982)

$$k_a = 4.1 \times 10^{-14} e^{-1440/T} cm^3 molecule^{-1} s^{-1}$$

= 3.3 x 10⁻¹⁶ cm³ molecule⁻¹ s⁻¹

at 298 K, with an estimated uncertainty of ±25% at 298 K;

$$k_b = 2.3 \times 10^{-12} e^{-2560/T} cm^3 molecule^{-1} s^{-1}$$

= 4.3 x 10⁻¹⁶ cm³ molecule⁻¹ s⁻¹

at 298 K with an estimated uncertainty of $\pm 30\%$ at 298 K. These expressions of Cowley et al. (1982) are recommended, with large uncertainties.

$$\frac{(CH_3)_3CO_2. + (CH_3)_3CO_2}{} + Products$$

Rate constants for the combination of t-butylperoxy radicals have been obtained by Parkes (1975) and Anastasi et al. (1978) using molecular modulation and flash photolysis techniques, respectively.

These data are in excellent agreement and, using an absorption cross-section for the t-butylperoxy radical of $4.0 \times 10^{-18} \text{ cm}^2$ at 240 nm (Anastasi et al., 1978), yield an observed rate constant at 298 K of

$$k_{obs} = (2.6 \pm 0.8) \times 10^{-17} \text{ cm}^3 \text{ molecule}^{-1} \text{ s}^{-1},$$

with an activation energy of ~8.8 kcal mol⁻¹ (Anastasi et al. 1978) over the narrow temperature range of 298-325 K. This temperature dependence is consistent with the Arrhenius expression obtained by Kirsch et al. (1978) of

$$k_{obs} = 1.7 \times 10^{-10} e^{-4775/T} cm^3 molecule^{-1} s^{-1}$$

Kirsch and Parkes (1981) have shown that the initial reaction leads predominantly to t-butoxy radicals, pathway (a), with pathway (b) being ~14% of the overall reaction at 298 K, decreasing rapidly with temperature.

$$(CH_3)_3CO_2$$
. + $(CH_3)_3CO_2$. + $2(CH_3)_3CO^*$ + O_2 (a)

$$+ (CH_3)_3COO(CH_3)_3 + O_2$$
 (b)

$\frac{\text{CH}_3\text{CO}_3 + \text{CH}_3\text{CO}_3 + \text{Products}}{2}$

This combination reaction has recently been studied by Addison et al. (1980) using molecular modulation absorption spectroscopy, and Basco and Parmar (1985) using a flash photolysis technique. The room temperature rate constants obtained disagree by a factor of ~3. While no recommendation can be made, it appears that this combination reaction is rapid.

In addition to these RO_2 self-combination reaction studies, rate constants for the reactions of $\mathrm{CH}_3\mathrm{O}_2$ with $(\mathrm{CH}_3)_3\mathrm{CO}_2$ and with $\mathrm{CH}_3\mathrm{C}(0)\mathrm{O}_2$ have been deduced (Parkes, 1975; Addison et al., 1980), with values at room temperature of

$$k[CH_3O_2 + (CH_3)_3CO_2] \sim 1 \times 10^{-13} \text{ cm}^3 \text{ molecule}^{-1} \text{ s}^{-1}$$

and

$$k[CH_3O_2 + CH_3C(O)O_2] = 3 \times 10^{-12} \text{ cm}^3 \text{ molecule}^{-1} \text{ s}^{-1}$$

Because of the inherent complexities of these experimental systems, no recommendations can be made.

In the absence of more experimental data, the rate constants and product ratios obtained for the ${\rm C_2H_5O_2}$, ${\rm (CH_3)_2CHO_2}$ and ${\rm (CH_3)_3CO_2}$ radicals are recommended as being reasonably representative of other primary, secondary and tertiary alkylperoxyradicals, and similarly with ${\rm CH_3CO_3}$ radicals. Furthermore, the use of a rate constant at 298 K of

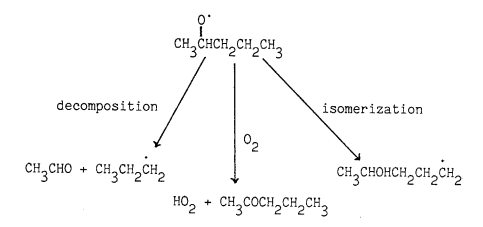
$$5 \times 10^{-12} \text{ cm}^3 \text{ molecule}^{-1} \text{ s}^{-1}$$

is recommended for the reaction of HO_2 radicals with all peroxy radicals (this being the major cross-combination reaction occurring).

Alkoxy Radical Reactions

It is now recognized that under atmospheric conditions the major alkoxy radical removal processes involve reaction with 0_2 , unimolecular decomposition and unimolecular isomerization (see, for example, Atkinson and Lloyd, 1984; Carter and Atkinson, 1985). These reactions are shown

for the case of the 2-pentoxy radical:



In addition, reactions with NO and NO_2 , though minor under most conditions, must be considered.

Reaction with 02

Absolute rate constants for the reactions of alkoxy radicals with 0_2 have been determined for CH $_3$ O (Sanders et al., 1980; Gutman et al., 1982; Lorenz et al., 1985; Wantuck et al., 1987), C_2H_5O (Gutman et al., 1982), and (CH $_3$) $_2$ CHO (Balla et al., 1985) radicals. The rate constants obtained are given in Table 7.

For the reaction of ${\rm CH_3O}$ radicals with ${\rm O_2}$, the rate constants of Gutman et al. (1982), Lorenz et al. (1985) and Wantuck et al. (1987) are in good agreement. These data are well fitted (Wantuck et al., 1987) by the double Arrhenius expression

$$k(CH_3O + O_2) = 1.5 \times 10^{-10} e^{-6028/T} + 3.6 \times 10^{-14} e^{-880/T}$$
 $cm^3 \text{ molecule}^{-1} \text{ s}^{-1}$

For the temperature range applicable to the troposphere, T $_{\rm z}$ 200-300 K, this expression reduces to

Table 7. Absolute Rate Constants for the Reactions of O_2 with Alkoxy Radicals

Radical	10^{15} x k $(\text{cm}^3 \text{ molecule}^{-1} \text{ s}^{-1})$	Т (К)	Reference
СH ₃ О	<2	295	Sanders et al. (1980)
	4.7 6.0 10.7 12.7	413 475 563 608	Gutman et al. (1982)
	1.9 ^a	298	Lorenz et al. (1985)
	2.3 ± 0.2 1.9 ± 0.2 4.0 ± 0.2 4.7 ± 0.3 4.4 ± 0.1 4.6 ± 0.2 7.2 ± 0.6 14 ± 1 12 ± 3 14 ± 1 30 ± 1 67 ± 4 140 ± 10 360 ± 110	298 298 348 423 423 423 498 573 573 573 673 773 873 973	Wantuck et al. (1987)
с ₂ н ₅ 0	8.0 9.8	296 353	Gutman et al. (1982)
(СН ₃) ₂ СНО	7.55 ± 0.30 7.06 ± 0.56 6.90 ± 0.37 8.22 ± 0.28 8.65 ± 0.26 7.48 ± 0.55 8.60 ± 0.36 9.16 ± 0.92 8.27 ± 0.80	294 295 296 296 314 330 346 367 384	Balla et al. (1985)

 $^{^{\}mathrm{a}}\mathrm{Calculated}$ from cited Arrhenius expression.

$$k(CH_3O + O_2) = 3.6 \times 10^{-14} e^{-880/T} cm^3 molecule^{-1} s^{-1}$$

which is similar to that of Lorenz et al. (1985) [obtained over the temperature range 298-450 K] of

$$k(CH_3O + O_2) = 5.5 \times 10^{-14} e^{-1000/T} cm^3 molecule^{-1} s^{-1}$$

This Arrhenius expression of Lorenz and Zellner (1985) is recommended [at 200 K, a 15% difference occurs between the rate constants calculated from the expressions of Lorenz et al. (1985) and Wantuck et al. (1987)].

For the ethoxy radical Gutman et al. (1982) obtained rate constants at 296 and 353 K. Combining these data of Gutman et al. (1982) with an estimated Arrhenius preexponential factor of 3.7 x 10^{-14} cm³ molecule⁻¹ s⁻¹ (from the preexponential factor for CH₃O + O₂, which yields 1.85 x 10^{-14} cm³ molecule⁻¹ s⁻¹ per abstractable H atom) leads to the expression for the ethoxy radical, and for primary alkoxy radicals in general, of

$$k(prim-alkoxy + O_2) = 3.7 \times 10^{-14} e^{-460/T} cm^3 3 molecule^{-1} s^{-1},$$

= 7.9 x 10⁻¹⁵ cm³ molecule⁻¹ s⁻¹ at 298 K.

Balla et al. (1985) determined absolute rate constants for the reaction of $\rm O_2$ with the $\rm (CH_3)_2CHO$ radical over the temperature range 294-384 K, and obtained the Arrhenius expression

$$k((CH_3)_2CHO + O_2) = 1.51 \times 10^{-14} e^{-(196 \pm 141)/T} cm^3 molecule^{-1} s^{-1}$$

Combining these rate constants of Balla et al. (1985) for the reaction of the isopropoxy radical with O_2 , and the estimated preexponential factor of $1.8 \times 10^{-14} \, \mathrm{cm}^3$ molecule⁻¹ s⁻¹, the following Arrhenius expression for secondary alkoxy radicals of

$$k(\text{sec-alkoxy} + 0_2) = 1.8 \times 10^{-14} \text{ e}^{-260/\text{T}} \text{ cm}^3 \text{ molecule}^{-1} \text{ s}^{-1},$$

= 7.5 x 10⁻¹⁵ cm³ molecule⁻¹ s⁻¹ at 298 K

is derived. These rate constant recommendations for primary and secondary alkoxy radicals are slightly different than those recommended by Carter and Atkinson (1985). These rate constants are expected to be applicable to substituted alkoxy radicals formed from, for example, the alkenes after initial OH radical reaction (for example, HOCH₂CH₂O).

Alkoxy Radical Decompositions

There have been a number of studies of these reactions in the gas phase. For the non-substituted alkoxy radicals; i.e., those formed from alkanes, the most useful of these experimental studies are those of Batt and co-workers (Batt, 1979a and references therein; Batt and Robinson, 1982) who determined the decomposition rates for a series of alkoxy radicals relative to their reaction rates with NO at ~400-470 K. Because of uncertainties concerning the absolute rates of the reference reactions and the relatively limited temperature range employed, the Arrhenius parameters, and thus the room temperature rate constants, for these reactions are subject to significant uncertainties.

As discussed by Carter and Atkinson (1985), while there have been several attempts to develop systematic schemes for estimating the

Arrhenius parameters for these reactions (Baldwin et al., 1977; Batt, 1979a; Choo and Benson, 1981), discrepancies have arisen primarily due to the differing thermochemical estimates used in the analyses of the experimental data. Batt and co-workers (Batt, 1979a) used the competition between the reactions

RO → decomposition

RO + NO → RONO

The rate constant for the RO + NO combination is derived for the decomposition rate constant data of RONO and an assumed zero temperature dependence for the RO + NO combination reaction. Carter and Atkinson (1985) used the alkoxy radical decomposition rates given by Batt (1979a) and Batt and Robinson (1982) since these data appeared to yield a reasonably self-consistent data set concerning the reactions of alkoxy radicals with O_2 , NO and NO₂. Batt (1979a) derived a value of ~4 x 10^{-11} ${\rm cm}^3$ molecule⁻¹ ${\rm s}^{-1}$ for the reaction of RO radicals with NO, while Choo and Benson (1981) recommended a value of $\sim 1 \times 10^{-11} \text{ cm}^3 \text{ molecule}^{-1} \text{ s}^{-1}$, both being independent of temperature. The recent study of Balla et al. (1985) on the kinetics of the reaction of $(CH_3)_2CHO$ radicals with NO yields a rate constant at ~400 K of ~(2.5-3.0) x 10^{-11} cm³ molecule⁻¹ s⁻¹, a factor of ~1.5 lower than estimated by Batt and co-workers (Batt, 1979a). Furthermore, a negative temperature dependence equivalent to an Arrhenius activation energy of -0.62 kcal mole -1 was determined by Balla et al. (1985). This leads to the preexponential factor for the RO + NOcombination reaction being lowered by a factor of 3 over that used by Batt (1979a), with a negative temperature dependence of \sim -0.6 kcal mole⁻¹. Thus,

$$k(RO + NO) = 1.3 \times 10^{-11} e^{300/T} cm^3 molecule^{-1} s^{-1}$$

This expression is consistent with the rate constant of $(2.08 \pm 0.12) \times 10^{-11} \text{ cm}^3 \text{ molecule}^{-1} \text{ s}^{-1}$ determined by Sanders et al. (1980) in the presence of 15 \pm 5 torr of SF₆ diluent (this reaction would still be in the fall-off regime under these conditions). Hence the Carter and Atkinson (1985) estimated preexponential factors for alkoxy radical decompositions will be lowered by a factor of 4, and the Arrhenius activation energies decreased by 0.6 kcal mole⁻¹.

The technique discussed by Carter and Atkinson (1985) involves the estimation of the Arrhenius preexponential factor, A, for the decompositions, and derivation of the Arrhenius activation energies, $E_{\rm decomp}, \ \mbox{by correlation with the heats of reaction } (\Delta H_{\rm decomp}). \ \mbox{The A}$ factors for decomposition to an alkyl radical and the corresponding carbonyl are now given by

$$A = n \times 2.7 \times 10^{14} \text{ s}^{-1}$$

where n is the reaction path degeneracy. In agreement with the earlier work of Baldwin et al. (1977) and Batt (1979a), an Evans-Polanyi plot of the Arrhenius activation energies, $E_{\rm decomp}$ (decreased by 0.6 kcal mole⁻¹ as discussed above), against the calculated heats of reaction, $\Delta H_{\rm decomp}$, yields a good straight line with no obvious effect of the leaving alkyl group, with

 $E_{\text{decomp}} = 11.4 + 0.70 \Delta H_{\text{decomp}}$

where E_{decomp} and ΔH_{decomp} are both in kcal mole⁻¹.

Table 8 summarizes the estimated heats of reaction and the calculated Arrhenius parameters and rate constants at 298 K for the classes of alkoxy radical decomposition which may be important in alkane photooxidations. The heats of reaction were estimated using the group additivity method of Benson (1976).

Dobé et al. (1986) have used a technique similar to that of Batt and co-workers to study the decomposition of the 2-pentoxy radical. Using the rate constant for the combination reaction of RO with NO given above, the data of Dobé et al. (1986) lead to a rate constant for 2-pentoxy decomposition to CH_3CHO and $\text{CH}_3\text{CH}_2\text{CH}_2$ of

$$k(decomposition) = 4.9 \times 10^{13} e^{-6700/T} s^{-1}$$

$$= 8.4 \times 10^3 \text{ s}^{-1} \text{ at } 298 \text{ K}$$

This is a factor of 1.9 higher than predicted from Table 8; considering the relatively long extrapolation from 363-418 K to 298 K this is reasonably good agreement.

Clearly, in view of the above discussion, these estimated alkoxy radical decomposition rate constants are subject to significant uncertainties, and direct determinations of these rate constants, obviating the need for reference reactions, are needed. It should also be noted that these decomposition reactions may be in the fall-off region between first-order and second-order kinetics at room temperature and

Table 8. Estimated Arrhenius Parameters and Room Temperature Rate Constants for Decompositions of Alkoxy Radicals

Reaction ^a	R.	ΔH _d ^b (kcal mole ⁻¹)	E _d (kcal mole ⁻¹)	k(298 K) ^c (s ⁻¹)
	СН3.	13.4	20.8	1.5 x 10 ⁻¹
0.	-CH ₂ .	9.1	17.8	2.4×10^{1}
O' RCH ₂ → R' + HCHO	>CH.	8.6	17.4	4.7×10^{1}
_	≯C.	6.9	16.2	3.5×10^2
	сн ₃ .	9.0	17.7	2.8×10^{1}
0 .	-CH ₂ .	4.7	14.7	4.5×10^3
RC-R' → R' + R'CHO	>CH-	4.2	14.3	8.8×10^3
O ' RC-R' + R' + R'CHO H	≯C.	2.5	13.2	5.6×10^4
	СН3.	6.6	16.0	5.0×10^2
0. 0	-СН ₂ .	2.3	13.0	7.9×10^4
0. 0 R-C-R' → R' + R'CR" R"	>CH·	1.8	.12.7	1.3×10^{5}
l R	≯C.	0.1	11.5	1.0 x 10 ⁶

^aThermochemical estimates based on R'-, R"- = -CH₃. Arrhenius parameters for R'-, R"- = CH₃-, >CH-, and \Rightarrow C- are assumed to be identical. bEstimated using group additivity and bond dissociation energies tabulated by Repsen (1076)

by Benson (1976). $^{\circ}$ Calculated using A = 2.7 x 10¹⁴ s⁻¹, and thus applicable only for a reaction path degeneracy of 1.

waddington, 1984). For the two alkoxy radicals for which pressure dependent decomposition rate constants have been observed [2-propoxy and 2-methyl-2-propoxy (t-butoxy)] the rate constants at room temperature and atmospheric pressure are apparently reasonably close to the limiting high pressure values (Batt and Robinson, 1982; Al Akeel and Waddington, 1984) [see also Table II in Baldwin et al. (1977), which predicts that for C₃ and higher alkoxy radicals, the corrections for fall-off behavior are small, being less than a factor of 2 at room temperature and atmospheric pressure]. Since the estimation procedure given above yields the limiting high pressure rate constants, the possibility that the reaction may not be at its high pressure and thus the true rate constants under atmospheric conditions may be lower should be recognized.

The above discussion has dealt with the reactions of alkoxy radicals formed from the alkanes, and the decompositions of other alkoxy radicals such as ${\rm HOCH_2CH_2\dot{O}}$ formed from ethene and ${\rm CH_3COCH(\dot{O})CH_3}$ formed from 2-butanone need to be considered.

For radicals of the structure $RCOCH(O)R^2$, the use of the above methods and equations used to estimate alkoxy radical deposition rates yields decomposition rates at the high pressure limit of:

for

$$\text{CH}_{3}\text{COCH}_{2}\text{O} + \text{CH}_{3}\text{CO} + \text{HCHO}$$

$$\text{k}_{298} = 1.0 \times 10^{4} \text{ s}^{-1}$$

for

$$\text{CH}_{3}\text{COCHCH}_{3} \rightarrow \text{CH}_{3}\text{CO} + \text{CH}_{3}\text{CHO}$$

$$\text{k}_{298} = 6.5 \times 10^{5} \text{ s}^{-1}$$

[these are somewhat uncertain due to uncertainties in the values of ΔH_{f} calculated for the alkoxy radicals (Benson, 1976)]. The reactions of these radicals with O_{2} at 760 torr total pressure and 298 K are estimated to have rates of 4.1 x 10^{4} s⁻¹ and 3.9 x 10^{4} s⁻¹, respectively. It is hence predicted for the $CH_{3}COCH(O)CH_{3}$ radical that decomposition will dominate over reaction with O_{2} by ~170:1. This prediction is in good agreement with experimental data (Batt, 1979b), and the estimation method outlined above appears to be applicable to these RCOCH(\dot{O})R' radicals.

These estimates of the rate constants for the reaction of alkoxy radicals with O_2 and for decomposition predict that for the reactions

$$CH_3CHC_2H_5 \rightarrow CH_3CH + C_2H_5$$
 (p)

 $k_{\rm b}/k_{\rm a}$ = 6.0 x 10¹⁷ cm³ molecule⁻¹ at 298 K. This rate constant ratio is a factor of approximately 4-5 lower than the direct measurements of $k_{\rm b}/k_{\rm a}$ of Carter et al. (1979b) and Cox et al. (1981) of 3.1 x 10¹⁸ molecule cm⁻³ at 303 K and 2.6 x 10¹⁸ molecule cm⁻³ at 296 K, respectively. In view of the fact that thermochemical estimates of heats of reaction, and hence of the estimated Arrhenius activation energies, are uncertain to at least ~1 kcal

 $mole^{-1}$, the minimum expected uncertainties in such rate constant ratio estimates are a factor of ~5 at 298 K.

Similarly, for radicals such as $\mathrm{CH_3OCH_2O}$ formed in the atmospheric degradation of dimethyl ether, the above estimation method predicts a decomposition rate at 298 K of 0.3 s⁻¹, in contrast to a rate of 4 x 10^4 s⁻¹ for reaction with $\mathrm{O_2}$. Hence reaction with $\mathrm{O_2}$ to yield $\mathrm{CH_3OCHO}$ should dominate, and this is experimentally observed (Tuazon, unpublished data).

For the alkoxy radicals $RR_1COHC(0)R_2R_3$ formed from the alkenes, however, estimations and experimental data are in serious disagreement (Baldwin et al., 1977; Batt, 1979a,c; Golden, 1979; Niki et al., 1978b). Thus, based upon the thermochemistries of the reactions

$$RR_1COHCR_2R_3 \rightarrow RR_1COH + R_2COR_3$$

and the estimation methods given above, the decompositions

and

have calculated rate constants at 298 K of ~10 s⁻¹ and 4.5 x 10^3 s⁻¹, respectively, compared to estimated rates for reaction with 0_2 (at 760 torr total pressure of air) of ~4 x 10^4 s⁻¹ for both radicals. Hence, it is predicted (as discussed previously by Baldwin et al., 1977; Golden, 1979; Batt, 1979a,c) that reaction with 0_2 will totally dominate for the ethene system and dominate for the 2-butenes.

However, experimentally, the RR₁COHC(\dot{o})R₂R₃ radicals decompose, with, at room temperature and 1 atmosphere or air, decomposition dominating for the \geq C₃ radicals, and with decomposition and reaction with O₂ being competitive for the HOCH₂CH₂O radical (Niki et al., 1978b, 1981b; Atkinson and Lloyd, 1984). Thus, for the HOCH₂CH₂O radical the rate constant ratio k_b/k_a for the reactions

$$HOCH_2CH_2O \rightarrow CH_2OH + HCHO$$
 (a)

$$HOCH_2CH_2\dot{O} + O_2 \rightarrow HOCH_2CHO + HO_2$$
 (b)

is $(5.4 \pm 1.0) \times 10^{-20} \text{ cm}^3 \text{ molecule}^{-1} \text{ s}^{-1} \text{ at room temperature } (~298 \text{ K})$ (Niki et al., 1981b). Using $k_b \sim 8 \times 10^{-15} \text{ cm}^3 \text{ molecule}^{-1} \text{ s}^{-1}$ (see above), this yields

$$k_a \sim 1.5 \times 10^5 \text{ s}^{-1}$$

at ~298 K and 700 torr total pressure. For the higher ($\geq C_3$) homologues of these radicals, the decomposition rates are expected to increase, as observed experimentally (Niki et al., 1978b; Atkinson et al., 1985). Thus, for these $\geq C_3$ homologues decomposition rates of $\geq 4 \times 10^5$ s⁻¹ are estimated.

Alkoxy Radical Isomerizations. No direct experimental data are available, but isomerization rate constants have been estimated initially by Carter (unpublished data, 1976) and subsequently, and in more detail, by Baldwin et al. (1977). The major relevant available experimental data concern measurements of the rate constant ratio for the reactions

$$_{\text{CH}}^{\text{$$

$$_{\text{CH}}^{2}\text{CH}^{2}\text{CH}^{2}\text{CH}^{3} + _{0}^{2} \rightarrow _{\text{HCCH}}^{2}\text{CH}^{2}\text{CH}^{3} + _{H0}^{2}$$
 (p)

from product yields determined in n-butane-NO $_{\rm x}$ -air (Carter et al., 1979b), HONO-n-butane-air (Cox et al., 1981) and n-butyl nitrite-air photolyses (Niki et al., 1981c). Rate constant ratios of $k_{\rm a}/k_{\rm b}$ of 1.65 x10¹⁹ molecule cm⁻³ at 303 K (Carter et al., 1979b), 1.9 x 10¹⁹ molecule cm⁻³ at 298 ± 2 K (Niki et al., 1981c) and 1.5 x 10¹⁹ molecule cm⁻³ at 296 K (Cox et al., 1981) were derived from these studies. These rate constant ratios are in good agreement, with an average value of $k_{\rm a}/k_{\rm b}=1.7$ x 10¹⁹ molecule cm⁻³ at ~299 K. Using the rate constant estimated as described above for $k_{\rm b}$, this leads to a rate constant of $k_{\rm a}=1.3$ x 10⁵ s⁻¹ at 299 K, a factor of ~4 lower than the estimate of Baldwin et al. (1977). Considering the large uncertainties in the estimation technique, this estimate is actually in fairly good agreement with the experimental data.

However, the data of Dóbé et al. (1986) for the isomerization of the 2-pentoxy radical lead to an isomerization rate of ~4 \times 10³ s⁻¹ at 298 K, significantly lower than the thermochemical estimates. The reasons for this discrepancy are not presently known, but may be due to the difficulties in quantitatively monitoring the end-products of this isomerization reaction (Dóbé et al., 1986).

As carried out by Carter and Atkinson (1985), the estimated Arrhenius parameters of Baldwin et al. (1977) have been modified to yield values of k_a which are a factor of 4 lower at 298 K, and the resulting Arrhenius parameters are given in Table 9 for 1,4-H shift and 1,5-H shift

Estimated Arrhenius Parameters and Room Temperature Rate Constants for 1,4- and 1,5-H Shift Isomerizations of Alkoxy Radicals Table 9.

Type of H-Shift	Type of Ring Strain ^a H-Shift (kcal mole ⁻¹)		Type of H E(Abstraction) ^b Abstracted (kcal mole ⁻¹)	$E_{\rm a}({\rm Isom})^{\rm c}$ (kcal mole ⁻¹)	A ^d (s ⁻¹)	k(298 K) (s ⁻¹)
1,4	5.9	-CH ₂ - >CH ₂ - >CH- -CH ₂ OH	7.7 4.6 6.5 3.4e	13.6 10.5 12.4 9.3	2.5 x 10 ¹¹ 1.7 x 10 ¹¹ 8.3 x 10 ¹⁰ 1.7 x 10 ¹¹ 8.3 x 10 ¹⁰	2.6 x 10 ¹ 3.4 x 10 ³ 1.7 x 10 ³ 1.4 x 10 ² 1.3 x 10 ⁴
ر تر	0.5	-сн ₃ -сн ₂ - >сн- -сн ₂ он -сн(он)-	7.7 4.6 4.6 6.5 3.4e	8.2 5.1 7.0 3.9	1.3 × 10 ¹¹ 8.4 × 10 ¹⁰ 4.2 × 10 ¹⁰ 8.4 × 10 ¹⁰ 4.2 × 10 ¹⁰	1.3 x 10 ⁵ 1.5 x 10 ⁷ 7.6 x 10 ⁶ 6.2 x 10 ⁵ 5.8 x 10 ⁷

bE(abstraction) = activation energy for abstraction by RO in bimolecular systems (i.e., no ring strain). Estimates of Baldwin et al. (1977), increased by 0.5 kcal mole, As estimated by Baldwin et al. (1977).

^oE_a(Isom) = E(abstraction) + ring strain.

dEstimates of Baldwin et al. (1977), decreased by a factor of 1.9 (see text), used.

eBaldwin et al. (1977) did not give an estimate for this abstraction. It is assumed that replacing -H with -OH decreases E(abstraction) by 1.2 kcal mole⁻¹, based on their estimates for abstraction from $-\mathrm{CH}_3$ and $-\mathrm{CH}_2\text{-groups}$. isomerizations of alkoxy radicals [the A factors of Baldwin et al. (1977) have been reduced by a factor of 1.9, and their estimated abstraction activation energies have been increased by 0.5 kcal $mole^{-1}$]. These estimates, however, must still be considered to be highly uncertain, and clearly further studies of these isomerization rate constants are needed.

While 1,2-, 1,3-, and 1, \geq 6-H shifts, via 3, 4 and \geq 7 member ring transition states, are also possible, the strain energies for the 3, 4 and 7-10 member rings are all higher than that estimated for the 1,4-H shift. Thus other H-atom shifts are expected to be much slower than 1,4-H shifts, which are themselves of marginal importance (Baldwin et al., 1977).

Reactions of RO Radicals with NO and NO_2 . Alkoxy radicals can also react with NO and NO_2 under atmospheric conditions

and

$$RO + NO_2 + RONO_2$$

+ $HNO_3 + R^O$

Absolute rate constants have been measured for the reactions of ${\rm CH_3O}$ and ${\rm (CH_3)_2CHO}$ radicals with NO and ${\rm NO_2}$, and the data are given in Table 10. The rate constants for the reactions of ${\rm CH_3O}$ with NO and ${\rm NO_2}$ measured by Sanders et al. (1980) and Lorenz et al. (1984) and by McCaulley et al. (1985), respectively, are in the fall-off region between second and third order kinetics (Gutman et al., 1982). The data of Lorenz et al. (1984) show that at 298 K the high pressure second order rate constant for the

Table 10. Absolute Rate Constants for Reaction of RO Radicals with NO and NO_2

Radical	Reactant	T (K)	P (torr)	10^{11} x k (cm ³ mole-cule ⁻¹ s ⁻¹)	Reference
сн30	NO	293	15 ± 5 (SF ₆)	2.08 ± 0.12	Sanders et al. (1980)
			9.8 (He)	1.1 ± 0.2	
		298	190 (He)	2.5 ± 0.5	Lorenz et al. (1984)
	NO ₂	250 298 390 473	4 (He) 4 (He) 4 (He) 4 (He)	0.531 ± 0.093 0.318 ± 0.049 0.120 ± 0.033 0.109 ± 0.027	McCaulley et al. (1985)
(CH ₃) ₂ CHO	NO	295 296 296 297 319 341 359 378	1 50 10 10 10 10 10	4.01 ± 0.21 3.31 ± 0.12 3.28 ± 0.07 3.54 ± 0.09 3.44 ± 0.07 3.10 ± 0.11 2.89 ± 0.10 2.68 ± 0.07	Balla et al. (1985)
	NO ₂	295 340 384	1-10 10 10	3.68 ± 0.16 2.59 ± 0.25 3.03 ± 0.15	Balla et al. (1985)

reaction of CH_3O with NO is $\geq 2.5 \times 10^{-11}$ cm³ molecule⁻¹ s⁻¹, with a low pressure bimolecular rate constant of ~4 x 10^{-12} cm³ molecule⁻¹ s⁻¹ at 298 K, the latter corresponding to the H-atom abstraction route. Similar data arise from the study of McCaulley et al. (1985) for the corresponding NO_2 reaction, from which the abstraction channel was concluded to have a rate constant of

k (CH₃0 + NO₂) = 9.6 x
$$10^{-12}$$
 e^{-1150/T} cm³ molecule⁻¹ s⁻¹
= 1.2 x 10^{-12} cm³ molecule⁻¹ s⁻¹ at 298 K.

For both addition reactions, negative temperature dependencies were observed (though these were in, or may have been in, the fall-off regions).

The kinetic data obtained by Balla et al. (1985) for the reactions of the $(CH_3)_2CHO$ radical with NO and NO_2 are at the high pressure limit, and show that these reactions have rate constants at room temperature of (3-4) \times 10^{-11} cm³ molecule⁻¹ s⁻¹, with, for the NO reaction, a slight negative temperature dependence.

A large amount of relative rate data have been obtained, and these are discussed in detail by Atkinson and Lloyd (1984). These relative rate data show that for the reaction of RO radicals with NO, the addition rate constants at ~400 K are ~3 x 10^{-11} cm³ molecule⁻¹ s⁻¹, with an uncertainty of \pm a factor of 3. For the CH₃O radical, this reaction has not reached the limiting high-pressure second-order limit at 700 torr total pressure of N₂, being ~20% below the high pressure limit at this pressure. For the RO + NO reactions, the H-atom abstraction process appears to be insignificant for R=CH₃, with k \leq 1.5 x 10^{-12} cm³ molecule⁻¹ s⁻¹ at

383-423 K (Batt and Rattray, 1979). Higher H-atom abstraction rate constants have been reported for the higher alkoxy radicals (see Atkinson and Lloyd, 1984; Morabito and Heicklen, 1985), with values of (3-6) x 10^{-12} cm³ molecule⁻¹ s⁻¹ at ~400 K being typical.

For the RO reactions with NO_2 , the relative rate data suggest that

$$k(RO + NO)/k(RO + NO_2) \sim 2 \pm 1$$

at ~400 K, and that the H-atom abstraction channel is negligible [with the most recent relative rate data yielding H-atom abstraction rate constants of ~1.5 x 10^{-13} cm³ molecule⁻¹ s⁻¹ for CH₃O and (CH₃)₂CHO radicals (Batt and Rattray, 1979; Batt, 1979b)].

The relative rate data are reasonably consistent with the few absolute rate constants available, and allow the following general recommendations for all RO radicals to be made:

$$k_{\infty}(RO + NO) = 1.3 \times 10^{-11} e^{300/T} cm^3 molecule^{-1} s^{-1}$$

= 3.6 x 10⁻¹¹ cm³ molecule⁻¹ s⁻¹ at 298 K,

with the H-atom abstraction rate constants being of minor importance at temperatures below 298 K.

$$k_{\infty}(RO + NO_2) \approx 1.3 \times 10^{-11} \text{ e}^{300/\text{T}} \text{ cm}^3 \text{ molecule}^{-1} \text{ s}^{-1}$$

 $\approx 3.6 \times 10^{-11} \text{ cm}^3 \text{ molecule}^{-1} \text{ s}^{-1} \text{ at } 298 \text{ K},$

with the H-atom abstraction process being of negligible importance at ≤ 298 K.

II. REACTIONS OF ALDEHYDES, KETONES AND α -DICARBONYLS

The aliphatic aldehydes and ketones are formed as intermediate "stable" chemical products during the atmospheric degradation reactions of a wide variety of organic compounds. These carbonyls arise from the reactions of the alkyl peroxy and alkoxy radicals discussed in Section I above. In addition, a number of α -dicarbonyls (for example, glyoxal, methylglyoxal and biacetyl) are formed as intermediate reactive products from the aromatic hydrocarbons (Darnall et al., 1979; Takagi et al., 1980; Atkinson et al., 1983b; Bandow and Washida, 1985a,b; Bandow et al., 1985; Tuazon et al., 1984a, 1986; Shepson et al., 1984; Dumdei and O'Brien, 1984; Gery et al., 1985). In the atmosphere, these carbonyls can photolyze and react with OH, NO3 and HO2 radicals, and these atmospheric reaction pathways are reviewed in this section. The corresponding atmospheric reactions of other classes of carbonyls such as the 1,4-unsaturated dicarbonyls are dealt with in Section XI.

Photolysis

In order to evaluate or calculate the photolysis rates of carbonyls under atmospheric and environmental chamber conditions, the radiation flux J, the carbonyl absorption cross-section, σ , and the photolytic quantum yield, ϕ , all as a function of wavelength, need to be known. Thus

$$k_{\text{photolysis}} = \int_{-290 \text{ nm}}^{-800 \text{ nm}} J_{\lambda} \sigma_{\lambda} \phi_{\lambda} d_{\lambda}$$

The radiation flux, J., is either experimentally measured or calculated for clear sky conditions and is not dealt with further here. The absorption cross-sections and quantum yields are obtained from experimental studies, and the available data are discussed below.

Formaldehyde. The absorption cross-sections and quantum yields have been most recently evaluated by NASA (DeMore et al., 1987) and IUPAC (Atkinson et al., 1988a). The most recent IUPAC recommendation (Atkinson et al., 1988a) accepts the absorption cross-section data of Moortgat et al. (1983). At longer wavelengths these absorption cross-sections are higher than those measured by Bass et al. (1980), but are confirmed by recent unpublished data of Biermann (University of California, Riverside). The recommended absorption cross-sections and quantum yields for the processes

$$HCHO + hv \rightarrow H + HCHO$$
 (a)

$$HCHO + hv \rightarrow H_2 + CO$$
 (b)

as recommended by the IUPAC panel (Atkinson et al., 1988a) are given in Table 11, although the higher wavelength resolution data of Moortgat et al. (1983 and references therein) should be used for modeling purposes (see Atkinson et al., 1988a).

Acetaldehyde. The quantum yields for the photolysis of acetaldehyde have been reviewed and evaluated by Atkinson and Lloyd (1984) and Baulch et al. (1984), based on the experimental studies of Horowitz and Calvert (1982), Horowitz et al. (1982) and Meyrahn et al. (1982). The recommended values as a function of wavelength are similar to within approximately 10%. The absorption cross-sections have also been reviewed by Baulch et

Table 11. Absorption Cross Sections, σ , and Quantum Yields, ϕ , for the Photolysis of HCHO (from Atkinson et al., 1988a)

λ (nm)	10 ²⁰ σ(cm ²) ^a 298 κ	^ф а	^ф b
270	0.95	0.38	0.43
280	1.80	0.57	0.32
290	2.93	0.73	0.24
300	4.06	0.78	0.21
310	4.60	0.78	0.22
320 ·	4.15	0.62	0.38
330	3.21	0.27	0.66
340	2.22	0	0.56 ^b
350	1.25	0	0.21 ^b
360	0.18	0	0.03 ^b

^aThe values are averaged for 10 nm intervals centered on the indicated wavelength. b760 torr total pressure of air.

al. (1982, 1984), and their recommendation is in good agreement with the recent values of Horowitz and Calvert (1982). The absorption cross-sections and quantum yields for the processes,

$$CH_3CHO + hv \rightarrow CH_4 + CO$$
 (a)

$$CH_3CHO + hv \rightarrow CH_3 + CHO$$
 (b)

$$CH_3CHO + hv \rightarrow CH_3CO + H$$
 (c)

as evaluated by Baulch et al. (1984), are given in Table 12. The pathway to form ketene plus $\rm H_2$ is negligible (Horowitz and Calvert, 1982; Horowitz et al., 1982).

<u>Propanal</u>. The sole data available for the photodissociation quantum yields for propanal arise from the studies of Shepson and Heicklen (1982a,b) and Heicklen et al. (1986). The more recent study of Heicklen et al. (1986) supercedes the previous work from the same laboratory. For the photodissociation

$$CH_3CH_2CHO + hv + C_2H_5 + CHO$$
 (a)

data were obtained at 294, 302, 313, 325 and 334 nm, with the quantum yields in air at 760 torr total pressure and 298 K being 0.89, 0.85, 0.50, 0.26 and 0.15 at these wavelengths, respectively. The quantum yield for the non-radical forming process

$$CH_3CH_2CHO + hv + C_2H_6 + CO$$
 (b)

Table 12. Absorption-Cross Sections, σ , and Quantum Yields ϕ_a and ϕ_b for CH₃CHO photolysis (at 1 atm air) [from Baulch et al., 1984]

Wavelength	10 ²⁰ σ/cm ²	^ф а	фb
260	2.0	0.46	0.31
270	3.4	0.31	0.39
280	4.5	0.05	0.58
290	4.9	0.01	0.53
295	4.5	0.0	0.48
300	4.3		0.43
305	3.4		0.37
315	2.1		0.17
320	1.8		0.10
325	1.1		0.04
330	0.69		0.0
335	0.38		
340	0.15		
345	0.08		

was observed to be essentially zero at wavelengths >313 nm. Absorption cross-sections have been presented by Calvert and Pitts (1966).

Butanal. The absorption cross-sections are as given by Calvert and Pitts (1966), and the gas-phase photolysis has been studied by Forgeteg et al. (1978, 1979) at 313 nm. At high pressure of added C_4F_8 , the quantum yields of the processes

$$C_3H_7CHO + hv \rightarrow C_3H_7 + CHO$$
 (a)

$$C_{3}H_{7}CHO + hv + C_{2}H_{4} + CH_{3}CHO$$
 (b)

were reported to be ϕ_b = 0.18, and ϕ_a ~0.3. Other photolysis pathways account for <5% of the overall quantum yield at 313 nm.

Higher Aldehydes. Absorption cross-section data are given by Calvert and Pitts (1966) for iso-butyraldehyde, and quantum yield data have been measured by Desai et al. (1986). At one atmosphere of air and room temperature the quantum yields for (CH₃)₂CH + HCO production were measured to be at the wavelengths: 253.7 nm, 0.20; 280.3 nm, 0.45; 302.2 nm, 0.55; 312.8 nm, 0.88; 326.1 nm, 0.88; and 334.1 nm, 0.69. For the higher aldehydes no quantum yield data are available, and further work is clearly needed concerning the photolysis of aldehydes (and ketones) under atmospheric conditions.

Acetone. The photodissociation of acetone has recently been studied under simulated atmospheric conditions by Gardner et al. (1984) and Meyrahn et al. (1986). Unfortunately, there are significant discrepancies between the results obtained from the studies of Gardner et al. (1984) and Meyrahn et al. (1986), with respect to both the quantum yields for radical

production and the effects of pressure on the quantum yields. At total pressures >350 torr of air, the quantum yield of the process

$$CH_3COCH_3 + hv \rightarrow CH_3CO + CH_3$$

was found to be 0.077 ± 0.0022 in the study of Gardner et al. (1984), independent of wavelength over the range 279-313 nm, and independent of temperature over the range 272-300 K.

Meyrahn et al. (1986) could not avoid the effects of secondary reactions, and added NO_2 to scavenge CH_3CO radicals to yield PAN ($CH_3C(0)OONO_2$). At 750 torr total pressure of air the quantum yields for CH_3CO formation were determined to be 0.55 at 280 nm, 0.30 at 290 nm, 0.15 at 300 nm, 0.05 at 310 nm, 0.028 at 320 nm, and ~0.033 at 330 nm. The absorption cross-section was also measured by Meyrahn et al. (1986). In addition, Cox et al. (1980) have reported an average photodissociation quantum yield of 0.33 \pm 0.06 for the wavelength region 280-330 nm, which is not consistent with the data of Gardner et al. (1984). The data of Meyrahn et al. (1986) are hence recommended, being more consistent with the variation of photodissociation quantum yields with wavelength observed for other ketones and with the data of Cox et al. (1980).

2-Butanone. Absorption cross-section data are given by Calvert and Pitts (1966). No definitive quantum yield data are available. By analogy with the quantum yields for the carbonyls discussed above, the photodissociation quantum yield is expected to be significantly less than unity. In fact, Carter et al. (1986) used a quantum yield for radical production of approximately 0.1 to fit environmental chamber irradiations of NO_X -2-butanone-air mixtures.

Higher Ketones. No data are available concerning the photodissociation quantum yields for the higher ketones. As for acetone, these quantum yields are likely to be low. However, since these compounds also react with the OH radical, it is expected that the OH radical reactions will be the dominant atmospheric loss process.

 α -Dicarbonyls. The α -dicarbonyls for which experimental data are available are glyoxal, methylglyoxal and biacetyl. The absorption crosssections have been measured and reported by Plum et al. (1983) over the wavelength regions 240-470 nm. In this wavelength region, two distinct absorption bands are observed, one from approximately 340-470 nm and a second from approximately 230-340 nm. Plum et al. (1983) studied the photolysis of these α -dicarbonyls in the longer wavelength band, although no wavelength resolution was carried out and only average photodissociation quantum yields could be derived. The α -dicarbonyl photolysis rates were determined relative to the measured photodissociation rate of ${\rm NO}_2$ in the environmental chamber employed. data obtained showed that photodissociation did occur in the longer wavelength band, and effective quantum yields (assuming a constant photodissociation quantum yield for wavelengths >325 nm) of 0.029 ± 0.018, 0.107 ± 0.030 and 0.158 ± 0.024 for glyoxal, methylglyoxal and biacetyl were derived.

For glyoxal, formaldehyde was observed as a product, showing that the process

 $(CHO)_2 + hv \rightarrow HCHO + CO$

must occur, with the HCHO yield corresponding to 13% of the glyoxal photolyzed (Plum et al., 1983). Other photodissociation pathways obviously occur. In the shorter wavelength band, Langford and Moore (1984) observed the formation of HCO radicals from the photolysis of glyoxal at 308 nm, with an estimated quantum yield of 0.4 \pm 0.2. Clearly, further work is necessary, but the available data suggest photodissociation into two HCO radicals with an appreciable quantum yield in the 230-340 nm absorption band, with photolysis to predominantly non-radical products [HCHO + CO and H $_2$ + 2 CO (Osamura et al. (1981)] occurring with low quantum yield in the first long wavelength band.

For methylglyoxal and biacetyl the photodissociation pathways are less well understood. However, the formation of PAN as a product of irradiated NO_{X} -methylglyoxal or biacetyl-air mixtures and the observation of the CH₃C(0)00 radical from the photolysis of biacetyl-0₂ mixtures (Cox et al., 1980) shows that the photolysis paths

$$CH_3COCHO + hv \rightarrow CH_3CO + CO$$

must occur. Indeed, Cox et al. (1980) have reported an average photodissociation quantum yield of 0.98 \pm 0.15 for biacetyl for the 280-330 nm wavelength region.

Hydroxyl Radical Reactions. The kinetics and mechanisms of the OH radical reactions with carbonyl compounds have been reviewed and evaluated by Atkinson (1986a), and updated evaluations for HCHO and CH_3CHO have been carried out for the IUPAC evaluation (Atkinson et al., 1988a). The

kinetic data for the simpler aldehydes and ketones, either as the recommended values or those measured but not recommended for lack of sufficient studies, are given in Table 13. All of these reactions proceed via overall H-atom abstraction, although it should be noted that for the aldehydes the initial reaction possibly involves OH radical addition to the >C=O bond system. Thus:

$$OH + HCHO \rightarrow H_2O + HCO$$

As noted in Section I, the HCO radical reacts rapidly with 0_2

$$HCO + O_2 \rightarrow HO_2 + CO$$

while the RCO radicals rapidly add 0_2 to form the corresponding acyl peroxy radicals (RC(0)00).

For glyoxal, Niki et al. (1985) have shown that the resulting HCOCO radical can either decompose or react with $\rm O_2$

$$HCOCO \rightarrow HCO + CO$$
 (a)

$$HCOCO + O_2 \rightarrow HCOC(0)OO$$
 (b)

$$\rightarrow$$
 2 CO + HO_2 (c)

Table 13. Room Temperature Rate Constants k and Arrhenius Parameters, $k = Ae^{-E/RT}$, for the Reaction of OH Radicals with Carbonyl Compounds. Except as Indicated, Taken from Atkinson (1986)

Carbonyl	10 ¹² k (298 K) cm ³ molecule ⁻¹ s ⁻¹	$10^{12} \times A$ cm^3 molecule ⁻¹ s ⁻¹	E/R (K)
ICHO ^a	11	16	110
:н ₃ сно ^а	16	5.6	-310
Н ₃ СН ₂ СНО	20		
H ₃ CH ₂ CH ₂ CHO	23		
СН ₃) ₂ СНСНО	27		
н ₃ (сн ₂) ₃ сно	27		
СН ₃) ₂ СНСН ₂ СНО	28		
сн ₃) ₃ ссно	27		
осн ₂ сно ^b	10		
н ₃ сосн ₃ а	0.23	1.7	600
н ₃ сосн ₂ сн ₃ с	1.1	2.3	170
H3COCH2CH2CH3	4.6		
H ₃ CH ₂ COCH ₂ CH ₃	1.8		
H ₃ COCH ₂ CH ₂ CH ₂ CH ₃	9.0		
н ₃ сн ₂ сосн ₂ сн ₂ сн ₃	6.8		
CHO) ₂	11		
н ₃ сосно	17		
H ₃ cococH ₃ d	2.3	1.1	450

aFrom Atkinson et al. (1988a). bFrom Niki et al. (1987a). cFrom Wallington and Kurylo (1987a). dFrom Dagaut et al. (1988b).

with $k_b \sim k_c$ and $k_a/k_b = 3.5 \times 10^{18}$ molecule cm⁻³. Thus, at 298 K and 760 torr total pressure of air, addition of 0_2 occurs 40% of the time, while formation of CO and $H0_2$ occurs the remaining 60% of the time. For methylglyoxal, the corresponding CH₃COCO radical is expected to decompose more rapidly, and hence 0_2 addition to yield CH₃COC(0)00 should be less important.

For the aldehydes and ketones, the positions of OH radical H-atom abstraction and the partial OH radical reaction rate constant at that position can be calculated using the estimation technique of Atkinson (1986b; 1987). As an example, the OH radical reaction with 2-butanone is calculated to proceed via the pathways

$$OH + CH_3CH_2COCH_3 \rightarrow H_2O + CH_2CH_2COCH_3$$
 (46%)

$$\rightarrow$$
 H₂O + CH₃CHCOCH₃ (46%)

$$+ \text{H}_2\text{O} + \text{CH}_3\text{CH}_2\text{COCH}_2$$
 (8%)

The subsequent reactions involve $\mathbf{0}_2$ addition to these radicals followed by the various reactions discussed above in Section I.

Ozone Reactions. For the carbonyls dealt with in this section which do not contain >C=C< bonds the reactions with O_3 are of negligible atmospheric importance, and indeed only upper limits to the rate constants of $<10^{-20}$ cm 3 molecule $^{-1}$ s $^{-1}$ have been obtained for HCHO, CH $_3$ CHO, (CHO) $_2$ and CH $_3$ COCHO (Atkinson and Carter, 1984).

 $\underline{\text{Nitrate Radical Reactions}}. \ \ \text{For the carbonyl compounds dealt with in} \\ \text{this section, rate constant data are available only for HCHO and CH$_3$CHO}$

(Atkinson et al., 1988a and references therein). Most of these studies are relative to the equilibrium constant for the reactions $NO_2 + NO_3 \ddagger N_2O_5$, and unfortunately, there are significant uncertainties in the value of this equilibrium constant (DeMore et al., 1987). For the reactions of the NO_3 radical with HCHO and CH_3CHO , the most recent NASA (DeMore et al., 1987) and IUPAC (Atkinson et al., 1988a) recommendations are in good agreement, with (DeMore et al., 1987; Atkinson et al., 1988a)

$$k(NO_3 + HCHO) = 6.0 \times 10^{-16} \text{ cm}^3 \text{ molecule}^{-1} \text{ s}^{-1} \text{ at 298 K}$$

and (Atkinson et al., 1988a)

$$k(NO_3 + CH_3CHO) = 1.4 \times 10^{-12} e^{-1860/T} cm^3 molecule^{-1} s^{-1}$$

over the range 264-374 K

=
$$2.7 \times 10^{-15} \text{ cm}^3 \text{ molecule}^{-1} \text{ s}^{-1} \text{ at } 298 \text{ K}.$$

With rate constants of this magnitude, these reactions with the ${\rm NO_3}$ radical are of minimal atmospheric importance as an aldehyde loss process, and the ${\rm NO_3}$ radical reaction rate constants for the ketones are expected to be lower, in the range of 10^{-17} to 10^{-16} cm³ molecule⁻¹ s⁻¹ at room temperature. However, these ${\rm NO_3}$ radical reactions with the aldehydes during nighttime hours can be important with respect to the products formed (Stockwell and Calvert, 1983; Cantrell et al., 1985, 1986). Thus, as with the corresponding OH radical reactions, these ${\rm NO_3}$ radical reactions proceed via H-atom abstraction

$$\sim 100_3 + RCHO \Rightarrow HNO_3 + RCO (R = H, alkyl)$$

and the production of HO_2 and $\mathrm{RC}(0)00$ radicals during nighttime hours can be significant.

 ${\rm HO_2}$ Radical Reactions. While the ${\rm HO_2}$ radical has been shown to react with HCHO (Su et al., 1979a,b; Niki et al., 1980; Veyret et al., 1982; Barnes et al., 1985), data are not available for other aldehydes. The room temperature rate constants obtained cover a range of a factor of 10, from 1 x ${\rm 10^{-14}~cm^3~molecule^{-1}~s^{-1}}$ (Su et al., 1979a,b) to 1.1 x ${\rm 10^{-13}~cm^3}$ molecule⁻¹ s⁻¹ (at 273 K) [Barnes et al., 1985]. The initial reaction involves addition of ${\rm HO_2}$,

$$HO_2$$
 + HCHO \rightarrow HOOCH₂O

followed by rapid isomerization of this intermediate species via a six-membered transition state

Su et al. (1979a,b) and Barnes et al. (1985) have proposed that the $00CH_2OH$ species can back-decompose to the HO_2 + HCHO reactants (presumably via the intermediacy of $HOOCH_2O$). That the $OOCH_2OH$ species is formed in this HO_2 radical reaction is shown by the formation, in the presence of NO_2 , of the peroxy nitrate $HOCH_2OONO_2$ (Niki et al., 1980). While further work is clearly necessary on this, and other analogous reaction(s), the data of Barnes et al. (1985) are recommended, i.e.,

 $k(HO_2 + HCHO + HOCH_2OO) = 1.1 \times 10^{-13} \text{ cm}^3 \text{ molecule}^{-1} \text{ s}^{-1} \text{ at 273 K}$ and

$$k(HOCH_2OO \rightarrow HO_2 + HCHO) = 20 s^{-1} at 273 K.$$

With this rapid back-decomposition rate of the ${\rm HOCH_200}$ radical, the reaction of ${\rm HO_2}$ radicals with HCHO is expected to be of minor importance.

III. ALKANES

The atmospheric chemistry of the alkanes has been reviewed and discussed in detail by Carter and Atkinson (1985), and the kinetics and mechanisms of the reactions with OH radicals and with $\rm O_3$ have recently been reviewed and evaluated (Atkinson and Carter, 1984; Atkinson, 1986a). The gas-phase reaction of the alkanes with $\rm O_3$ are of negligible importance as an atmospheric loss process, since the available data show that the rate constants for these reactions at room temperature are $\rm <10^{-23}~cm^3$ molecule $\rm ^{-1}~s^{-1}$ (Atkinson and Carter, 1984). Under atmospheric conditions, the potential loss processes for the alkanes involve gas-phase reactions with OH and $\rm NO_3$ radicals.

OH Radical Reactions

The kinetics and mechanisms of the reactions of the OH radical with alkanes have been critically reviewed and evaluated by Atkinson (1986a). Rate constants have been determined over significant temperature ranges for a number of alkanes, and it is evident, as expected from theoretical considerations, that the Arrhenius plots are curved. Accordingly, the three-parameter expression

$$k = C T^2 e^{-D/T}$$

has been used. The recommended 298 K rate constants and the parameters C and D given by Atkinson (1986a) are shown in Table 14 for alkanes of relevance to tropospheric chemistry. Room temperature rate constants for other alkanes for which recommendations were not given (generally due to only single studies being carried out) are also given in Table 14.

These OH radical reactions proceed via H-atom abstraction from the C-H bonds and, as discussed by Atkinson (1986a,b), the rate constants for these OH radical reactions with alkanes can be fit to within a factor of 2 over the temperature range 250-1000 K by consideration of the $\rm CH_3$ -, $\rm -CH_2$ - and $\rm >CH$ - groups in the alkane and the substituent groups around these $\rm CH_3$ -, $\rm -CH_2$ - and $\rm >CH$ - groups. Thus

$$k(CH_3-X) = k_{prim}^{O} F(X)$$

$$k(X-CH_2-Y) = k_{sec}^{o} F(X) F(Y)$$

and

$$k(X-CH \stackrel{Y}{\underset{Z}{\sim}}) = k_{tert}^{O} F(X) F(Y) F(Z)$$

where k_{prim}^{0} , k_{sec}^{0} and k_{tert}^{0} are the OH radical rate constants per -CH₃, -CH₂- and >CH-group, respectively, and F(X), F(Y) and F(Z) are the substituent factors for X, Y and Z substituent groups. As derived by Atkinson (1986b, 1987)

Table 14. Rate Constants at 298 K and Parameters C and D in $k=CT^2e^{-D/T}$ for the Reaction of OH Radicals with Alkanes (from Atkinson, 1986a)

Alkane	10 ¹² x k (298 K) cm ³ molecule ⁻¹ s ⁻¹	10^{18} x C cm ³ molecule ⁻¹ s ⁻¹	D (K)
Methane	0.00841	6.95	1280
Ethane	0.274	13.7	444
Propane	1.18	12.7	-14
n-Butane	2.53	a	a
2-Methylpropane	2.37	9.58	-305
n-Pentane	4.04		
2-Methylbutane	3.9		
2,2-Dimethylpropane	0.852	17.5	-179
n-Hexane	5.58		
2-Methylpentane	5.5		
3-Methylpentane	5.6		
2,2-Dimethylbutane	2.6		•
2,3-Dimethylbutane	6.2	b	ь р
n-Heptane	7.2		
2,4-Dimethylpentane	5.1		
2,2,3-Trimethylbutane	4.1		
n-Octane	8.72	c	С
2,2,4-Trimethylbutane	3.66	d	d
2,2,3,3-Tetramethylbutane	1.06	18.7	-133
n-Nonane	10.0		
n-Decane	11.2		
n-Undecane	13.3		

Table 14 (continued) - 2

Alkane	$10^{12} \text{ x k } (298 \text{ K})$ cm ³ molecule ⁻¹ s ⁻¹	$10^{18} \times C$ cm ³ molecule ⁻¹ s ⁻¹	D (K)
n-Dodecane	13.9		
n-Tridecane	15.5		
Cyclopropane	0.07		
Cyclobutane	1.2		
Cyclopentane	5.2		
Cyclohexane	7.38	е	е
Cycloheptane	13.1		
Methylcyclohexane	10.3		

^aArrhenius expression of $k = 1.55 \times 10^{-11} \, e^{-540/T} \, cm^3 \, molecule^{-1} \, s^{-1}$ recommended (300-500 K).

^bIndependent of temperature over the range ~300-500 K.

^cArrhenius expression of $k = 3.12 \times 10^{-11} \, e^{-380/T} \, cm^3 \, molecule^{-1} \, s^{-1}$

recommended (300-500 K).

dArrhenius expression of $k = 1.62 \times 10^{-11} e^{-443} cm^3 molecule^{-1} s^{-1}$ recommended (300-500 K).

 $e_{\text{Arrhenius expression of k}} = 2.73 \times 10^{-11} e^{-390/\text{T}} \text{ cm}^3 \text{ molecule}^{-1} \text{ s}^{-1}$ recommended (300-500 K).

$$k_{\text{prim}}^{0} = 4.47 \times 10^{-18} \text{ T}^{2} \text{ e}^{-303/\text{T}} \text{ cm}^{3} \text{ molecule}^{-1} \text{ s}^{-1},$$
 $k_{\text{sec}}^{0} = 4.32 \times 10^{-18} \text{ T}^{2} \text{ e}^{233/\text{T}} \text{ cm}^{3} \text{ molecule}^{-1} \text{ s}^{-1},$
 $k_{\text{tert}}^{0} = 1.89 \times 10^{-18} \text{ T}^{2} \text{ e}^{711/\text{T}} \text{ cm}^{3} \text{ molecule}^{-1} \text{ s}^{-1},$
 $F(-\text{CH}_{3}) = 1.00$

$$F(-\text{CH}_{2}^{-}) = F(>\text{CH}_{-}) = F(>\text{C}<) = e^{76/\text{T}} = 1.29 \text{ at } 298 \text{ K}.$$

This estimation technique not only allows the calculation of OH radical reaction rate constants for alkanes for which experimental data do not exist, but also allows the isomeric alkyl radical distribution to be calculated for a given alkane. Thus, for example, for 2-methylbutane the calculated OH radical reaction rate constants at 298 K at the various carbon atoms are (in units of 10^{-12} cm³ molecule⁻¹ s⁻¹)

The percentages of OH radical reaction calculated to occur at each of the carbons are also given. These alkyl radicals then react as discussed above in Sections I and II, and the expected subsequent reaction sequences in the presence of NO are as follows (where RONO₂ is the corresponding alkyl nitrate formed from the alkyl peroxy radicals).

$$(CH_3)_2 CHCHCH_3 + O_2 \longrightarrow (CH_3)_2 CHCHCH_3$$

$$\downarrow NO \\ RONO_2$$

$$(CH_3)_2 CHCHCH_3 + NO_2$$

$$\downarrow CH_3 CHCHCH_3 + NO_2$$

$$\downarrow O_2, NO \\ \downarrow O_2, NO \\ RONO_2$$

$$(CH_3)_2 CHO + NO_2$$

$$\downarrow O_2$$

$$CH_3 COCH_3 + HO_2$$

$$(CH_3)_2CHCH_2CH_2 + O_2 \rightarrow (CH_3)_2CHCH_2CH_2OO$$

$$\downarrow NO \\ CH_3)_2CHCH_2CH_2O^2 + NO_2$$

$$\downarrow O_2$$

$$(CH_3)_2CHCH_2CHO + HO_2$$

In the absence of NO, the alkyl peroxy radicals will react with ${\rm HO}_2$ and ${\rm RO}_2$ radicals.

For the longer chain (>C $_3$) alkanes, alkoxy radical isomerization can also occur in addition to decomposition and reaction with O $_2$ (Section I and Carter and Atkinson, 1985). For example, for the 2-pentoxy radical

$$\begin{array}{c} \text{CH}_3\text{CHCH}_2\text{CH}_2\text{CH}_3 & \xrightarrow{\text{isom}} & \text{CH}_3\text{CHOHCH}_2\text{CH}_2\text{CH}_2 \\ & \downarrow \text{O}_2, & \text{NO} \\ & & \downarrow \text{RONO}_2 \\ \\ \text{CH}_3\text{CHOHCH}_2\text{CH}_2\text{CH}_2\text{O} \\ & \downarrow \text{isom} \\ \\ \text{CH}_3\text{COHCH}_2\text{CH}_2\text{CH}_2\text{OH} \\ & \downarrow \text{O}_2 \\ \\ \text{CH}_3\text{C(O)CH}_2\text{CH}_2\text{CH}_2\text{OH} \\ \end{array}$$

It should be noted, however, that this reaction sequence has not been experimentally confirmed, and the fraction of the reaction of the δ-hydroxyalkyl peroxy radical reacting with NO to yield the corresponding δ-hydroxyalkyl nitrate has not been experimentally determined. The limited data available concerning alkyl nitrate formation from these hydroxy-substituted alkyl peroxy radicals (from computer model fits to environmental chamber data) suggest that this alkyl nitrate formation is minimal, and Carter and Atkinson (1985) recommend that this fractional alkyl nitrate formation from the reaction

$$\begin{array}{c} \text{OO.} \\ \text{RCHOHCH}_2\text{CH}_2\text{CHR.} + \text{NO} \rightarrow \text{RCHOHCH}_2\text{CH}_2\text{CH(ONO}_2)\text{R.} \end{array}$$

is essentially zero.

NO3 Radical Reactions

It has been shown that the ${\rm NO_3}$ radical reacts with the alkanes, with room temperature rate constants being in the 10^{-17} to 10^{-16} cm³ molecule⁻¹ s⁻¹ range (Atkinson et al., 1988b). These kinetic data were obtained using a relative rate technique. Using rate constants for the reactions of ${\rm NO_3}$ radicals with n-heptane and 2,3-dimethylbutane of 1.36 x 10^{-16} and 4.06 x 10^{-16} cm³ molecule⁻¹ s⁻¹ at 296 K, respectively (Atkinson et al., 1988b), the rate constants given in Table 15 are derived. Under atmospheric conditions, the nighttime reactions of the alkanes with the ${\rm NO_3}$ radical can be calculated to be typically a factor of 100 less important as an atmospheric loss process compared to the daytime OH radical reaction.

These ${\rm NO}_3$ radical reactions, similar to the OH radical reactions, proceed via H-atom abstraction from the C-H bonds

$$NO_3 + RH \rightarrow HNO_3 + R$$

followed by the alkyl radical reactions. The distribution of alkyl radical isomers formed from the more complex alkanes are not known, but the reactivities of primary, secondary and tertiary C-H bonds are anticipated to be tertiary >> secondary >> primary.

Table 15. Rate Constants for the Reaction of NO $_3$ Radicals with Alkanes at 296 \pm 2 K (Taken from Atkinson et al., 1988b)

Alkane	10^{16} x k $(\text{cm}^3 \text{ molecule}^{-1} \text{ s}^{-1})^a$
n-Butane	0.6
n-Pentane	0.8
n-Hexane	1.1
n-Heptane	1.4
n-Octane	1.8
n-Nonane	2.4
2-Methylpropane	1.0
2,3-Dimethylbutane	4.1
Cyclohexane	1.3

 $^{^{\}mathrm{a}}$ Uncertainty limits are of the order of $\pm 15-35\%$.

IV. HALOALKANES

The kinetics and mechanisms of the OH radical and O_3 reactions with the haloalkanes have recently been reviewed and evaluated by Atkinson (1986a) and Atkinson and Carter (1984), respectively. The gas-phase reactions of the haloalkanes with O_3 are of negligible importance as an atmospheric loss process, since the available data show that the rate constants for these reactions are $<10^{-20}$ cm³ molecule-1 s-1 at room temperature (Atkinson and Carter, 1984). The potential tropospheric loss processes for the haloalkanes then involve reaction with OH and NO_3 radicals and photolysis. To date, no data exist for the NO_3 radical reactions, but by analogy with the alkanes it is expected that these NO_3 radical reactions will be slow (with rate constants $<10^{-17}$ cm³ molecule-1 s-1 at room temperature for the C_1 and C_2 haloalkanes at room temperature).

OH Radical Reactions

The kinetics and mechanisms of the reactions of the OH radical with the haloalkanes have been critically reviewed and evaluated by Atkinson (1986a). Rate constants have been determined over significant temperature ranges for a number of haloalkanes, and the Arrhenius plots are curved, as expected from theoretical considerations. Atkinson (1986a) accordingly evaluated the data in terms of the modified expression

$$k = C T^2 e^{-D/T}$$

The recommended 298 K rate constants and the parameters C and D given by Atkinson (1986a) are given in Table 16 for haloalkanes of tropospheric interest, together with the room temperature rate constants for other haloalkanes for which recommendations were not made.

For the haloalkanes containing F, Cl and Br substituents, these OH radical reactions proceed by H atom abstraction, and in the absence of experimental data the estimation technique of Atkinson (1986a,b, 1987) can be used to calculate the distribution of haloalkyl radicals formed. Garraway and Donovan (1979) have reported that CF_3I and other non-hydrogen-containing iodine containing haloalkanes such as C_2F_5I and C_3F_7I react with OH radicals, presumably by I atom abstraction to form HOI and the corresponding haloalkyl radical. The atmospheric reactions of the haloalkyl radicals have been dealt with in Section I above.

Photolysis

Photolysis is a potentially important tropospheric loss process for the haloalkanes, especially for those containing multiple Cl and/or Br atoms and no H atoms (thus precluding OH radical reaction). Absorption cross-sections for ${\rm CCl_4}$, ${\rm CFCl_3}$, ${\rm CF_2Cl_2}$, ${\rm CHF_2Cl}$, ${\rm CH_3Cl}$ and ${\rm CH_3CCl_3}$ are tabulated in the NASA evaluation (DeMore et al., 1985, 1987). Absorption cross-sections for other haloalkanes have been measured by Chou et al (1978), Robbins (1977), Hubrich and Stuhl (1980), Hubrich et al. (1977), Vanlaethem-Meuree et al. (1978), Green and Wayne (1976/77) and Molina et al. (1982). The absorption spectra are unstructured and continuous, and the photodissociation quantum yields are expected, consistent with experimental observations for ${\rm CCl_4}$, ${\rm CFCl_3}$ and ${\rm CF_2Cl_2}$ (Baulch et al.,

Table 16. Room Temperature Rate Constants and Temperature Dependencies $(k=CT^2e^{-D/T})$ for the Gas-Phase Reactions of the OH Radical with Haloalkanes (from Atkinson, 1986a)

Haloalkane	$10^{14} \text{ x k}(298 \text{ K})$ cm ³ molecule ⁻¹ s ⁻¹	$10^{18} \times C$ cm ³ molecule ⁻¹ s ⁻¹	D (K)
CH ₃ Cl	4.4	3.5	585
CH ₃ Br	3.9	1.2	296
CH ₂ FC1	4.4	3.8	604
CH ₂ Cl ₂	14	8.5	500
CHF ₂ C1	0.47	1.5	1000
CHFC12	3.0	1.7	479
CHC13	10	6.3	504
CF ₃ Br	<0.1		
CF ₂ Cl ₂	<0.04		
CF ₂ ClBr	<0.1		
CFC13	<0.05	•	
CC1 ₄	<0.4		
CH3CH2Cl	40		
CH ₃ CHF ₂	3.4		
CH2ClCH2Cl	22		
CH ₂ BrCH ₂ Br	25		
CH3CF2C1	0.36	2.1	1171
CH3CC13	1.2	5.9	1129
CH ₂ FCF ₃	0.85	1.3	769
CH ₂ ClCF ₂ Cl	~1.5	•	
CHFC1CF ₃	1.0	0.91	624
CHC1 ₂ CF ₃	3.4	а	a

 $a_{k} = 1.2 \times 10^{-12} e^{-1056/T} cm^{3} molecule^{-1} s^{-1}$ over the range 245-375 K.

1982), to be unity, and at shorter wavelengths (<230 nm) two halogen atoms can be simultaneously in the primary process. The halogen atom produced is a Br atom for bromine-containing haloalkanes and a Cl atom for haloalkanes containing Cl atoms but no Br atoms.

$$CF_2ClBr + hv \rightarrow CF_2Cl + Br$$

$$CF_2Cl_2 + hv + CF_2Cl + Cl$$

The subsequent reactions of the haloalkyl radicals formed have been discussed in Section I above.

In general, photolysis of the fluorochloroalkanes is insignificant in the troposphere. Indeed, photolysis of CF_3Br is extremely slow in the troposphere, with a calculated lifetime of >1000 yr (Molina et al., 1982). Only for haloalkanes containing one Br and one Cl atom (for example, CF_2ClBr) or two Br atoms (such as CF_2Br_2) or more does photolysis in the troposphere become important (Molina et al., 1982).

V. ALKENES

As discussed by Atkinson and Lloyd (1984), Atkinson and Carter (1984) and Atkinson (1986a), the atmospheric loss processes of the alkenes are by reaction with OH and NO_3 radicals and O_3 . The conjugated dialkenes also react with NO_2 , and this reaction could be of some importance in environmental chamber experiments carried out at NO_2 concentrations higher than ambient.

OH Radical Reactions

The kinetics and mechanisms of the reactions of the OH radical with the alkenes, cycloalkenes and dialkenes have been reviewed and evaluated by Atkinson (1986a). For ethene and the methyl-substituted ethenes the OH radical reaction proceeds essentially totally by OH radical addition to the carbon-carbon double bond, with H atom abstraction from the side chains accounting for <5% of the total reaction for propene and 2methylpropene (Hoyermann and Sievert, 1979, 1983) at room temperature and <10% for cis- and trans-2-butene (Hoyermann and Sievert, 1983). For 1butene, the recent data of Hoyermann and Sievert (1983) and Atkinson et al. (1985) show that H atom abstraction accounts for <10% of the overall reaction at room temperature. To date, only for 1,3- and 1,4cyclohexadiene has H atom abstraction been shown to occur to any significance (Ohta, 1984), with this process accounting for 8.9% and 15.3%of the overall OH radical reactions with 1,3- and 1,4-cyclohexadiene, respectively, at room temperature. However, for the alkenes with side chains, a small amount of H atom abstraction must occur, with this process contributing only a small fraction of the overall reaction. Assuming that H atom abstraction from substituent alkyl groups occurs at the same rate as in the alkanes (Atkinson, 1987), then this H atom abstraction pathway is generally insignificant. For an extreme case, H atom abstraction in 1heptene is then calculated to account for 10-15% of the overall OH radical reaction.

The rate constants for these O'! radical reactions with the alkenes are at, or are very close to, the high pressure second-order kinetic limit at atmospheric pressure. For ethene, this high pressure limit appears to

be attained at approximately 760 torr of air diluent gas. Thus, the high pressure second order rate constants will be reasonably applicable throughout the troposphere. The room temperature rate constants and temperature dependent parameters for the reactions of the OH radical with a series of atmospherically important alkenes, cycloalkenes and dialkenes are given in Table 17.

As discussed above, except for 1,3- and 1,4-cyclohexadiene (and presumably also the corresponding cycloheptadienes), OH radical addition to the >C=C< bond is the dominant reaction pathway. The OH radical can add to either carbon atom of the double bond(s), and Cvetanovic (1976) has reported that for propene addition to the terminal carbon occurs 65% of the time

OH +
$$CH_3CH=CH_2$$
 + CH_3CHCH_2 and CH_3CHCH_2OH (~35%) (~65%)

The β -hydroxyalkyl radicals then rapidly add O_2 under atmospheric conditions, as discussed in Section I above. In the presence of NO, these β -hydroxyalkyl peroxy radicals form NO_2 plus the corresponding β -hydroxyalkoxy radical, with a small amount of β -hydroxyalkyl nitrate also being formed (Shepson et al., 1985)

Table 17. Rate Constants k at 298 K and Arrhenius Parameters ($k = Ae^{-E/RT}$) for the Reaction of OH Radicals with Alkanes at the High Pressure Limit (from Atkinson, 1986a)

Alkene	10 ¹² x k(298 K) (cm ³ molecule ⁻¹ s ⁻¹)	$10^{12} \times A$ (cm ³ molecule ⁻¹ s ⁻¹)	E/R(K)
Ethene	8.54	2.15	-411
Propene	26.3	4.85	-504
1-Butene	31.4	6.53	-468
1-Pentene	31.4		
3-Methyl-1-butene	31.8	5.32	- 533
1-Hexene	37		
3,3-Dimethyl-1-butene	28.4	•	
1-Heptene	40		
2-Methylpropene	51.4	9.51	-503
2-Methyl-1-butene	60.7		
cis-2-Butene	56.1	10.9	-488
cis-2-Pentene	65.1		
<u>trans</u> -2-Butene	63.7	10.1	-549
trans-2-Pentene	67		
2-Methyl-2-butene	86.9	19.2	- 450
2,3-Dimethyl-2-butene	110	: •	
Cyclopentene	67.0		
Cyclohexene	67.4		,
Cycloheptene	74.1		
1-Methylcyclohexene	95		

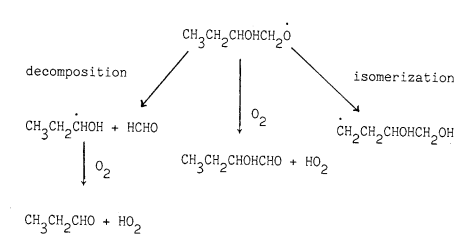
(continued)

Table 17 (continued) - 2

Alkene	$10^{12} \times k(298 \text{ K})$ (cm ³ molecule ⁻¹ s ⁻¹)	$10^{12} \times A$ (cm ³ molecule ⁻¹ s ⁻¹)	E/R(K)
α-Pinene	53.2	12.0	-444
β-Pinene	78.2	23.6	-357
Δ^3 -Carene	87.0		
d-Limonene	169		
1,3-Butadiene	66.8	13.9	-468
2-Methyl-1,3-butadiene	101	25.5	-409

At atmospheric pressure and room temperature, Shepson et al. (1985) determined yields of $\text{CH}_3\text{CHOHCH}_2\text{ONO}_2$ and $\text{CH}_3\text{CH}(\text{ONO}_2)\text{CH}_2\text{OH}$ of approximately 0.016 for each nitrate.

The β -hydroxyalkoxy radicals can then decompose, react with O_2 or isomerize, as discussed in Section I above. Thus, for the alkoxy radical formed after internal addition of the OH radical to 1-butene:



The experimental data for the simpler alkenes show that, at room temperature, under atmospheric conditions decomposition dominates over reaction with O_2 . Indeed, for the $\ge C_3$ alkanes the reaction with O_2 is negligible and only the products arising from decomposition are observed at room temperature and atmospheric pressure. For ethene, Niki et al. (1981b) have shown that both reaction with O_2 and decomposition of the HOCH₂CH₂O radical occurs

$$HOCH_2CH_2O \rightarrow CH_2OH + HCHO$$
 (a)

$$HOCH_2CH_2O + O_2 + HOCH_2CHO + HO_2$$
 (b)

with $k_b/k_a = (5.4 \pm 1.0) \times 10^{-20}$ at 298 K.

For the $\mathrm{CH_3CH_2CHOHCH_2\dot{O}}$ radical formed from 1-butene, the experimental data of Atkinson et al. (1985) show that isomerization is not important. This observation is in accord with the estimates of Atkinson and Lloyd (1984).

Hence, apart from ethene, for which reaction of the $\mathrm{HOCH_2CH_2O}$ radical with $\mathrm{O_2}$ and decomposition are competitive at 298 K and atmospheric pressure, the β -hydroxyalkoxy radicals formed subsequent to OH radical reactions with the simpler alkenes decompose. For propene the reaction scheme in the presence of NO is then

Similar reaction schemes have been expected to apply to the conjugated dialkenes. For example, for isoprene (2-methyl-1,3-butadiene) the reaction sequence is expected to be (Lloyd et al., 1983) [with the reactions at the two double bonds being dealt with separately, and neglecting nitrate formation (initial stable products are underlined)]:

OH +
$$CH_2$$
= $CHC(CH_3)$ = CH_2 + CH_2 = $CHC(CH_3)$ CH₂OH and CH_2 = $CHC(CH_3)$ CH₂

NO \downarrow NO₂

NO \downarrow NO₂
 CH_2 = $CHC(CH_3)$ CH₂OH

 CH_2 CH

 CH_2 CH

The products formed depend on which double bond OH radical addition occurs at, but not at which carbon atom of the particular double bond. For conjugated dialkenes, the estimation technique of Ohta (1983) allows the fraction of the overall OH radical addition reaction proceeding at each >C=C< double bond to be calculated [note that this information cannot be obtained from the estimation technique of Atkinson (1986a)]. Thus, for isoprene, OH radical addition to the CH₂=CH- and CH₂=C< bonds

OH +
$$CH_2$$
= $CHC(CH_3)$ = CH_2 + $HOCH_2$ - $CHC(CH_3)$ = CH_2 and CH_2 CHC(CH_3)= CH_2 (a)

$$\circ$$
H \circ CH₂=CHC(CH₃)CH₂ and CH₂=CHC(CH₃)CH₂OH (b)

is calculated to be in the ratio k_a/k_b = 34/66. The reactions of the products have been dealt with in Section II above (carbonyls) or in Section VIII below (α,β -unsaturated carbonyls).

However, two recent studies of the products formed from the gas-phase reaction of the OH radical with isoprene show that methyl vinyl ketone and methacrolein (together with their co-product HCHO) do not account for the entire reaction pathway. Thus, Gu et al. (1985) observed that at room temperature and atmospheric pressure of air or O_2 the major products were methyl vinyl ketone, methacrolein and 3-methylfuran in approximate respective yields of 16%, 23% and 5%. More recently, Tuazon et al. (unpublished data, 1988) have observed methyl vinyl ketone, methacrolein and HCHO as products, with yields of ~30%; ~20% and ~60%, respectively, with the HCHO yield being in reasonable agreement with the sum of the methyl vinyl ketone and methacrolein yields. Additionally, 3-methylfuran is a product of

this reaction (with a yield of 4.4%). These data of Tuazon et al. show that -40% of the overall reaction pathways are not presently accounted for by the reaction scheme presented by Lloyd et al. (1983) [see above], and IR absorption bands due to other products were indeed observed.

The production of 3-methylfuran presumably occurs by one of two routes

and initial OH radical addition at the other >C=C< bond will also yield 3-methylfuran by these schemes.

03 Reaction

The kinetics and mechanisms of the gas-phase reactions of 0_3 with the alkanes, cycloalkenes and dialkenes have been reviewed and evaluated by Atkinson and Carter (1984). The kinetic data for alkenes of atmospheric importance [updated to take into account the more recent studies of Bahta et al. (1984) and Bennett et al. (1987)] are given in Table 18. These reactions proceed by initial 0_3 addition to the >C=C< bond to yield an energy-rich ozonide which rapidly decomposes to a carbonyl and an initially energy-rich biradical.

$$O_{3} + R_{2} C - C < R_{3} + R_{2} C - C < R_{4} + R_{2} C - C < R_{4} + R_{2} C - C < R_{4}$$

$$C - C < R_{4} + R_{2} C - C < R_{4}$$

$$C - C < R_{4} + R_{2} C - C < R_{4}$$

$$C - C < R_{4} + R_{2} C - C < R_{4}$$

$$C - C < R_{3} + R_{4} C - C < R_{4}$$

$$C - C < R_{4} + R_{5} C - C < R_{5} + R_{5} C - C < R_{5}$$

$$C - C < R_{5} + R_{5} C - C < R_{5} + R_{5} C - C < R_{5}$$

$$C - C < R_{5} + R_{5} C - C < R_{5} + R_{5} C - C < R_{5}$$

$$C - C < R_{5} + R_{5} C - C < R_{5} + R_{5} C - C < R_{5}$$

$$C - C < R_{5} + R_{5} C - C < R_{5} + R_{5} C - C < R_{5} + R_{5} C - C < R_{5}$$

$$C - C < R_{5} + R_{5} C - C < R_{5$$

where $[\]^{\ddagger}$ denotes an energy-rich species. Based upon the data of Herron and Huie (1978), it is assumed that $k_a \sim k_b$ for the alkene systems. The energy-rich biradicals can then be collisionally stabilized or unimolecularly decompose.

$$[R_1 R_2 \dot{COO}]^{\ddagger} + M \rightarrow R_1 R_2 \dot{COO} + M$$

$$[R_1R_2COO]^{\ddagger}$$
 + products.

Table 18. Rate Constants k at 298 K and Arrhenius Parameters ($k = Ae^{-E/RT}$) for the Gas-Phase Reactions of O_3 with Alkenes, Cycloalkenes and Dialkenes (from Atkinson and Carter, 1984, except as indicated)

Alkene	$10^{18} \text{ x k}(298 \text{ K})$ (cm ³ molecule ⁻¹ s ⁻¹)	$10^{15} \times A$ (cm ³ molecule ⁻¹ s ⁻¹)	E/R (K)
Ethene ^a	1.7	12	2630
Propene	11.3	13.2	2105
1-Butene	11.0	3.46	1713
1-Pentene	10.7		
1-Hexene	11.7		
1-Heptene	17.3		
2-Methyl-2-propene	12.1	3.55	1693
<u>cis</u> -2-Butene	130	3.52	983
<u>cis</u> -2-Pentene	209		
<u>trans</u> -2-Butene	200	9.08	1136
trans-2-Pentene	315		
2-Methyl-2-butene	423	6.17	798
2,3-Dimethyl-2-butene	1160	3.71	347
Cyclopenteneb	275-813		
Cyclohexeneb	104-160		
Cycloheptene	319		
α-Pinene	84	0.96	731
β-Pinene	21		
Δ ³ -Carene	120		
d-Limonene	640		
1,3-Butadiene ^c	7.5	26	2430
2-Methyl-1,3-butadiene	14.3	12.3	2013

^aFrom Atkinson et al. (1988a). ^bIncludes the data of Bennett et al. (1987). ^cIncludes the data of Bahta et al. (1984).

The fraction of the initially formed biradical which is collisionally stabilized is thus expected to be pressure dependent, and this has been confirmed by the study of Hatakeyama et al. (1984), in which the stabilized biradicals formed from a variety of alkenes were "trapped" by reaction with $S0_2$ to form sulfate. For the \underline{trans} -2-butene system, the fraction of the biradical stabilized was a function of the total pressure of air, increasing from essentially zero at zero total pressure to a highpressure limit of 0.185 attained at ~600 torr total pressure of air. At one atmosphere total pressure, the fractional amount of stabilized biradicals formed from ethene, propene, trans-2-butene and 2-methyl-2propene were 0.390, 0.254, 0.185 and 0.174, respectively (Hatakeyama et al., 1984). This yield of stabilized biradicals from ethene is in excellent agreement with previous values of 0.38 (Su et al., 1980), 0.37 (Kan et al., 1981) and 0.35 (Niki et al., 1981d), and that for $\underline{\text{trans}}$ -2butene agrees well with the stabilized biradical yield of 0.18 obtained by Niki et al. (1977) from cis-2-butene.

Thus, at ~760 torr total pressure of air and ~298 K the fractions of $[CH_200]^{\ddagger}$ and $[CH_3CH00]^{\ddagger}$ stabilized from the ethene and 2-butene systems are 0.37 and 0.18, respectively. Assuming that $k_a = k_b$ for the decomposition routes of the initially formed ozonide and that the $[CH_200]^{\ddagger}$ and $[CH_3CH00]^{\ddagger}$ biradicals formed from propene react identically to those formed from ethene and the 2-butenes, these stabilization yields predict a total stabilized biradical yield from propene (CH_200) plus $CH_3CH00)$ of 0.275, in good agreement with the observed yield of 0.254 \pm 0.023 (Hatakeyama et al., 1984).

However, the data of Hatakeyama et al. (1984) for the stabilized biradical yield from 2-methyl-2-propene (0.174) are not consistent with

the above data for the stabilization yield of $[CH_2OO]^{\ddagger}$ (0.37) and the stabilization yield of ~0.30 for the $[(CH_3)_2COO]^{\ddagger}$ biradical in the 2,3-dimethyl-2-butene system (Niki et al., 1987b). This suggests that the stabilization yields of these biradical species, as may be expected, are dependent on the reaction system in which they are formed.

Data concerning the decomposition pathways of the energy-rich biradicals are available mainly for $[CH_200]^{\ddagger}$, $[CH_3CH00]^{\ddagger}$ and $[(CH_3)_2C00]^{\ddagger}$ (Atkinson and Lloyd, 1984; Atkinson and Carter, 1984; Niki et al., 1987b). For the $[CH_200]^{\ddagger}$ biradical, formed from the reaction of O_3 with ethene, the recommendation of Atkinson and Lloyd (1984) yields (at ~760 torr total pressure of air and 298 K):

$$[\dot{c}H_{2}0\dot{o}]^{\ddagger} \xrightarrow{M} \dot{c}H_{2}0\dot{o}$$

$$\rightarrow co_{2} + H_{2}$$

$$\rightarrow co + H_{2}0$$

$$\rightarrow H + Hco_{2}$$

$$H + co_{2}$$

Thus, for the reaction of 0_3 with ethene at 760 torr total pressure of air and room temperature, the overall reaction stoichiometry is

$$0_3$$
 + CH_2 = CH_2 + HCHO + 0.37 CH_2 00 + 0.12 $H0_2$ + 0.13 H_2 + 0.19 $C0_2$ + 0.44 CO + 0.44 H_2 0

For the [CH₃CHOO][‡] biradical, a fraction of 0.18 is stabilized at 298 K and atmospheric pressure. Based upon the recommendations of Atkinson and Lloyd (1984) for the decomposition routes, the reactions of this biradical under atmospheric conditions is given by

$$[CH3CHOO]^{\ddagger} \rightarrow CH3CHOO$$
 (18%)

$$+ CH_3 + CO + OH$$
 (~30%)

$$+ CH_3 + CO_2 + H$$
 (~32%)

$$+$$
 HCO $+$ CH₃O (~6%)

$$+ CH_{4} + CO_{2}$$
 (14%)

Thus, when combined with reactions of the $[CH_2OO]^{\ddagger}$ biradical, the overall stoichiometry of the reaction of O_3 with propene at ~760 torr total pressure and 298 K is then expected to be

$$^{0}_{3}$$
 + $^{CH}_{3}$ CH= $^{CH}_{2}$ + 0.50 HCHO + 0.50 CH $_{3}$ CHO + 0.185 CH $_{2}$ OO + 0.09 CH $_{3}$ CHOO + 0.25 HO $_{2}$ + 0.15 OH + 0.03 CH $_{3}$ O + 0.31 CH $_{3}$ O $_{2}$ + 0.07 CH $_{4}$ + 0.065 H $_{2}$ + 0.325 CO $_{2}$ + 0.40 CO + 0.22 H $_{2}$ O

However, Carter et al. (1986) observed from computer model simulations of propene-NO $_{\rm X}$ -air irradiations that the radical formation from this scheme is too high. Clearly further study of the products arising from the reaction of O $_{\rm X}$ with propene under atmospheric pressure are needed.

The stabilized biradicals are known to react with aldehydes, ${\rm SO_2}$, ${\rm CO}$, ${\rm H_2O}$, and ${\rm NO_2}$, and it is expected that they will also react with NO.

RCHOO + R'CHO + RCH
$$\stackrel{\circ}{}_{0}$$
 CHR'

RCHOO + SO₂ $\stackrel{H_2O}{\longrightarrow}$ RCHO + H_2 SO₄

RCHOO + CO + products

RCHOO + H_2O + RCOOH + H_2O

RCHOO + H_2O + RCHO + H_2O

Based upon the available data for the reactions of CH_2OO radicals with these reactants [with rate constants relative to the reaction of CH_2OO with SO_2 of: HCHO, ~0.25 (Su et al., 1980); CO, 0.0175 (Su et al., 1980); H_2O , (2.3 ± 1) x 10^{-4} (Suto et al., 1985) and NO_2 , 0.014 (Manzanares et al., 1985)], it appears that the reaction of stabilized biradicals with water vapor will be their dominant loss process, leading to the formation of carboxylic acids.

Recently Niki et al. (1987b) have studied the products and mechanism of the reaction of 0_3 with 2,3-dimethyl-2-butene under atmospheric conditions. A fraction of 0.25-0.30 of the initially energy-rich biradical $[(CH_3)_2COO]^{\ddagger}$ was observed to be stabilized at atmospheric pressure, with the major decomposition route of this biradical involving the isomerization

$$[(CH_3)_2COO]^{\ddagger} \rightarrow [CH_3C(OOH)=CH_2]$$

followed by dissociation of this unsaturated hydroperoxide intermediate

$$[CH_3C(OOH)=CH_2] \rightarrow CH_3COCH_2 + OH$$

The above discussion shows that only for the reaction of O_3 with ethene (and maybe 2,3-dimethyl-2-butene) are the products reasonably well known. Even for propene there are significant uncertainties in the radicals formed and their yields under atmospheric conditions.

NO3 Radical Reaction

The rate constants for the gas-phase reactions of the NO₃ radical with a large number of alkenes, cycloalkenes and dialkenes have been determined using absolute and relative rate methods. A reliable absolute rate constant at room temperature is available for <u>trans-2-butene</u> (Ravishankara and Mauldin, 1985; Dlugokeneky and Howard, 1988), and this enables the recent relative rate constant data of Atkinson et al. (1988b) to be placed on an absolute basis. The room temperature rate constants for a series of atmospherically important alkenes are given in Table 19.

Table 19. Room Temperature Rate Constants k for the Reaction of NO₃ Radicals with a Series of Alkenes (from Atkinson et al., 1988b)

Alkene	$k (cm^3 molecule^{-1} s^{-1})$
Ethene	2.1 x 10 ⁻¹⁶
Propene	9.4×10^{-15}
1-Butene	1.2×10^{-14}
2-Methylpropene	3.1×10^{-13}
<u>cis</u> -2-Butene	3.5×10^{-13}
trans-2-Butene	3.9×10^{-13}
2-Methyl-2-butene	9.3×10^{-12}
2,3-Dimethyl-2-butene	5.7×10^{-11}
Cyclopentene	4.6×10^{-13}
Cyclohexene	5.3×10^{-13}
Cycloheptene	4.8×10^{-13}
α-Pinene	5.8×10^{-12}
ß-Pinene	2.4×10^{-12}
Δ ³ -Carene	1.0×10^{-11}
d-Limonene	1.3×10^{-11}
1,3-Butadiene	9.8×10^{-14}
2-Methyl-1,3-Butadiene	5.9 x 10 ⁻¹³

These reactions proceed by NO $_3$ radical addition to the >C=C< bond, with H atom abstraction being totally insignificant,

followed by rapid addition of 0_2

$$\begin{array}{c}
R_1 & ONO_2 \\
\downarrow & C \\
R_2
\end{array}$$

$$+ O_2 \longrightarrow R_4 & OOO. R_4$$

When ${\rm NO_3}$ radicals are present, NO concentrations are extremely low, and these ${\rm g-nitratoalkyl}$ peroxy radicals will then either react with ${\rm HO_2}$ and other ${\rm RO_2}$ radicals, or reversibly add ${\rm NO_2}$ to yield the thermally unstable nitrato peroxynitrates

Because of the expected rapid thermal decomposition of the peroxy nitrates, these species act as a temporary reservoir of the β -nitratoalkyl peroxy radicals. Of course, if NO is present at sufficiently high concentrations to react with the peroxy radicals, then the corresponding alkoxy radical will be formed (this radical can also be formed from the

 $RO_2 + RO_2$ radical reactions)

These alkoxy radicals can then react with O_2 , decompose or isomerize, as discussed in Section I above. For the major $\mathrm{B-nitratoalkoxy}$ radical formed from the reaction of NO_3 radicals with propene (by NO_3 radical addition at the terminal carbon atom), Shepson et al. (1985) showed that decomposition and reaction with O_2 both occur

$$CH_3CH(0)CH_2ONO_2 \rightarrow CH_3CHO + HCHO + NO_2$$

$$CH_3CH(0)CH_2ONO_2 + O_2 \rightarrow CH_3COCH_2ONO_2 + HO_2$$

NO₂ Reactions

 ${
m NO}_2$ has been shown to react with conjugated dialkenes with rate constants at room temperature of >10^{-20} {
m cm}^3 molecule⁻¹ s⁻¹ (Atkinson et al., 1984b; Ohta et al., 1986). These reactions are of negligible importance as an atmospheric loss process of the alkenes and dialkenes, but may be marginally important in environmental chamber experiments carried out at high ${
m NO}_2$ concentrations. The articles of Atkinson et al. (1984), Niki et al. (1986) and Ohta et al. (1986) should be consulted for further details of the kinetics and mechanisms of these reactions.

VI. HALOALKENES

The kinetics and mechanisms of the gas-phase reactions of the haloalkenes with $\rm O_3$ and OH radicals have been reviewed and evaluated by Atkinson and Carter (1984) and Atkinson (1986a), respectively. The loss processes which are potentially important in the troposphere involve reaction with OH and NO $_3$ radicals and with O $_3$.

OH Radical Reactions

The kinetics and mechanisms of the gas-phase reactions of the haloalkanes with the OH radical have been reviewed by Atkinson (1986a). More recent kinetic and product data have been reported by Edney et al. (1986a,b), Tuazon et al. (1988) and Winer et al. (1987). Room temperature rate constants and temperature dependent parameters for the atmospherically important haloalkenes are given in Table 20. These reactions proceed entirely, or essentially entirely, by OH radical addition to the >C=C< bond, and are at the second-order high pressure limit at atmospheric pressure. Indeed, since the limiting high-pressure rate constant for CH2=CHCl is attained at a total pressure of argon of <50 torr, these reactions will be at the high pressure second-order limit throughout the troposphere. The OH radical adds preferentially to the least halogen substituted carbon atom of the >C=C< bond, and the resulting substituted alkyl radical will rapidly add O_2 under atmospheric conditions:

Table 20. Rate Constants k at Room Temperature and Arrhenius Parameters ($k = Ae^{-E/RT}$) for the Reaction of OH Radicals with Haloalkenes

Haloalkene	10^{12} x k (cm ³ molecule ⁻¹ s ⁻¹)	$10^{12} \times A \cdot (cm^3 \text{ molecule}^{-1} \text{ s}^{-1})$	E/R(K)
CH ₂ =CHCl ^a	6.60	1.48	- 390
CH2=CCl2p	8.11		
eis-CHCl-CHClb	2.38	0.563	-427
trans-CHCl=CHClb	1.80		
CHCl=CCl ₂ a	2.36		
CCl ₂ =CCl ₂ ^a	0.167	9.64	1209
CH ₂ =CHCH ₂ Cl ^e	17		
<u>eis</u> -CH ₂ ClCH=CHCl ^b	8.45		
trans-CH ₂ ClCH=CHC1 ^b	14.4		

^aFrom Atkinson (1986a). ^bFrom Tuazon et al. (1988). ^cFrom Edney et al. (1986a) and Winer et al. (1987).

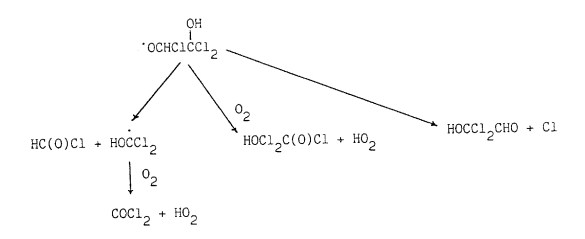
OH + CHCl=CCl₂ + HOCHClCCl₂ and CHClCCl₂ (minor)
$$\downarrow o_2 \qquad \qquad \downarrow o_2$$
HOCHClCCl₂ · OOCHClCCl₃

In the absence of sufficient NO, these peroxy radicals will react with ${\rm HO}_2$ and other ${\rm RO}_2$ radicals. In the presence of sufficiently high concentrations of NO, reaction with NO will occur to yield ${\rm NO}_2$ and the corresponding alkoxy radical.

The subsequent reactions of these hydroxy-haloalkoxy radicals are not totally understood (Tuazon et al., 1988); reaction with $\rm O_2$, decomposition or Cl atom elimination are possible:

HOCHC1CC1₂

$$\downarrow O_2$$



The available data (Edney et al., 1986a,b; Tuazon et al., 1988) show that Cl atom elimination occurs during the OH radical reactions with all of the chloroethenes except for CH_2 =CHCl, and this process competes with other decomposition processes of the alkoxy radical. For vinyl chloride, a unit yield of HCHO and HC(0)Cl are observed (Tuazon et al., 1988). For the remaining chloroethenes, the fraction of the overall alkoxy radical proceeding by C-C bond decomposition is approximately as follows: cis-1,2-dichloroethene, ~0.3; trans-1,2-dichloroethene, ~0.35; 1,1-dichloroethene, ~0.7-0.75; trichloroethene, ~0.4; and tetrachloroethene, ~0.25, with the remainder of the overall reactions occurring by O_2 reaction with and/or Cl atom elimination from the alkoxy radicals.

For allyl chloride, the observed products are CH_2CICHO and HCHO in similar yield (~25%), together with smaller, and similar, amounts of $HOCH_2CHO$ and HC(O)CI (Winer et al. 1987). Edney et al. (1986a) observed a further variety of products, including $CH_2CICOCH_2CI$ which shows that CI atoms were involved in secondary reactions in their system. Addition of OH radicals to the >C=C< bond is expected to be the major (~98%) initial reaction pathway, leading (in the presence of NO) to the alkoxy radicals $OCH_2CHOHCH_2CI$ and $HOCH_2CH(O)CH_2CI$. Decomposition of these lead to the products observed by Winer et al. (1987)

OCH₂CHCH₂Cl
$$\rightarrow$$
 HCHO $+$ CH₂ClCHOH
$$\downarrow o_2$$

$$CH_2ClCHO + HO_2$$

and

$$HOCH_2CHCH_2C1 \rightarrow CH_2C1CHO + CH_2OH$$

$$\downarrow O_2$$

$$HCHO + HO_2$$

$$\uparrow O_2$$

$$\uparrow O_2$$

$$\uparrow O_2$$

$$\downarrow O_2$$

$$\downarrow O_2$$

$$\uparrow O_2$$

$$\downarrow O_2$$

$$\downarrow O_2$$

$$\uparrow O_2$$

$$\downarrow O_2$$

$$\downarrow O_2$$

Reaction of the 'OCH₂CHOHCH₂Cl radical with O₂ to yield CH₂ClCHOHCHO may occur, accounting, at least in part, for the other product(s) not observed by Winer et al. (1987). Photolysis of HC(0)Cl in the irradiated NO_x-air system employed by Edney et al. (1986a) to produce Cl atoms with subsequent chain reactions involving Cl atoms may explain many of the other products observed by Edney et al. (1986a). The products of the reaction of OH radicals with cis- and trans-1,3-dichloropropenes in the presence of NO were HC(0)Cl and CH₂ClCHO in essentially unit yield, irrespective of the 1,3-dichloropropene isomer. This observation shows that the alkoxy radicals CH₂ClCHOHCHClO and CH₂ClCH(0)CHClOH must both decompose with essentially unit efficiency, similar to the \geq C₃ alkenes.

03 Reaction

The kinetics and mechanisms of the gas-phase reactions of 0_3 with the haloalkenes have been reviewed and evaluated by Atkinson and Carter (1984), and data have since been reported for allyl chloride [CH2=CHCH2Cl] (Edney et al., 1986a; Winer et al., 1987). The measured rate constants at room temperature for the reaction of 0_3 with atmospherically important haloalkenes are given in Table 21. No temperature dependencies have been measured to date. The reactivities of the haloalkenes with respect to reaction with 0_3 are much lower than the analogous alkenes, and these 0_3 reactions are of minor or negligible importance as a haloalkenes loss process in the troposphere. As for the alkenes, these reactions proceed by 0_3 addition to the >C=C< bond, followed by rapid decomposition of this energy-rich ozonide:

$$CH_{2}=CHC1 + O_{3} \longrightarrow \begin{bmatrix} O & O & 0 \\ CH_{2} & CHC1 \end{bmatrix}^{\ddagger}$$

$$A & b$$

$$CH_{2}=CHC1 + O_{3} \longrightarrow \begin{bmatrix} CH_{2}OO]^{\ddagger} + HC(O)C1 \end{bmatrix}$$

In contrast to the alkene reactions, for which it is generally assumed that $k_a \sim k_b$, Zhang et al. (1983) have determined that $k_a/k_b \sim 3.2$ for the decomposition of the ozonide formed from this reaction of 0_3 with vinyl chloride, and that the fraction of the initially energy rich $[CH_200]^{\ddagger}$ biradical which is stabilized at atmospheric pressure is 0.25. This lower fraction of stabilization of the $[CH_200]^{\ddagger}$ biradical formed from vinyl

Table 21. Room Temperature Rate Constants k for the Reactions of 0_3 with a Series of Haloalkenes (from Atkinson and Carter, 1984, except as indicated)

Haloalkene	$k (cm^3 molecule^{-1} s^{-1})$
CH ₂ =CHCl	2.4 x 10 ⁻¹⁹
CH ₂ =CCl ₂	3.7×10^{-21}
cis-CHCl=CHCl	$<5 \times 10^{-21}$
trans-CHCl=CHCl	1.2 x 10 ⁻¹⁹
CHCl=CCl ₂	<3 x 10 ⁻²⁰
CCl ₂ =CCl ₂	<2 x 10 ⁻²³
CH ₂ =CHCH ₂ Cl ^a	1.6×10^{-18}
cis-CHCl=CHCH ₂ Cl	1.5 x 10 ⁻¹⁹
trans-CHCl=CHCH ₂ Cl	6.7 x 10 ⁻¹⁹

 $^{^{\}mathrm{a}}$ From Edney et al. (1986a) and Winer et al. (1987).

chloride than from ethene was rationalized by the fact that $[CH_2^{00}]^{\ddagger}$ formed from vinyl chloride is ~10 kcal mol⁻¹ more energized than $[CH_2^{00}]^{\ddagger}$ formed from ethene.

From a study of the reaction of 0_3 with $\underline{\text{trans}}$ -1,2-dichloroethene, Niki et al. (1983a, 1984) have shown that the $[HC(C1)00]^{\ddagger}$ biradical reacts via

$$[HC(C1)00]^{\ddagger} \xrightarrow{M} ... (\sim 23\%)$$

$$\rightarrow HC(O)C1 + O(^{3}P) (\sim 20\%)$$

$$\rightarrow C1 + H + CO_{2}$$

$$\rightarrow C1 + OH + CO$$

$$\rightarrow other products (\sim 57\%)$$

Furthermore, it was proposed that the stabilized HC(Cl)00 radical reacts with water (homogeneously or heterogeneously) to yield peroxyformic acid

$$HC(C1)00 + H_20 + HC(0)00H + HC1$$

Peroxyformic acid yields of 0.04, 0.09, 0.23 and 0.10 were observed from the reactions of 0_3 with CH₂=CHCl, cis-CHCl=CHCl, trans-CHCl=CHCl and CH₂=CCl₂, respectively (Niki et al., 1982b).

The major products from the reactions of 0_3 with <u>cis-</u> and <u>trans-1,3-</u> dichloropropene were determined to be HC(0)Cl (initial yield ~0.7), CH₂ClCHO (yield ~0.4) and CH₂ClCOOH (Tuazon et al., 1984b). These products can be formed by the general schemes shown above for the alkenes and the haloalkenes.

NO3 Radical Reaction

Rate constants have been determined at room temperature for the reaction of NO_3 radicals with the chloroethenes and allyl chloride (Table 22). These kinetic data indicate that these NO_3 radical reactions will be a minor loss process for the haloalkenes under tropospheric conditions. These reactions are expected to proceed by NO_3 radical addition to the >C=C< bond, totally analogous to the NO_3 radical reactions with the alkenes. Although no product or mechanistic data are available, the subsequent reactions are expected to be analogous to those for the alkenes, with the added complication that C1 atom elimination may occur.

Table 22. Room Temperature Rate Constants k for the Reactions of $^{\rm NO}_3$ Radicals with Haloalkenes (from Atkinson et al., 1988b)

Haloalkene	$10^{16} \text{ x k (cm}^3 \text{ molecule}^{-1} \text{ s}^{-1})$
CH ₂ =CHCl	4.5
CH ₂ =CCl ₂	13
<u>eis</u> -CHCl=CHCl	1.5
trans-CHCl=CHCl	1.1
CHCl=CCl ₂	2.9
CCl ₂ =CCl ₂	<0.62
CH ₂ =CHCH ₂ Cl	5.6

VII. ALKYNES

The potentially important reactions of the alkynes are with OH and NO_3 radicals and O_3 . At the present time, for the simpler alkynes kinetic and product data are available for acetylene, propyne, 1-butyne, and 2-butyne.

OH Radical Reaction

The limiting high-pressure second-order rate constants at room temperature are given in Table 23. For acetylene, the rate constant is in the fall-off region between second- and third-order kinetics below ~1000 torr (Schmidt et al., 1985; Wahner and Zetzsch, 1985). At 298 K and 760 torr total pressure of air, Atkinson (1986a) recommended that k(acetylene) = $7.8 \times 10^{-13} \text{ cm}^3 \text{ molecule}^{-1} \text{ s}^{-1}$, with an Arrhenius activation energy of 0.46 kcal mol⁻¹. At 298 K, the rate constant for acetylene is given by

$$k = \left(\frac{k_0[M]}{1 + \frac{k_0[M]}{k_{\infty}}}\right) 0.6 \left\{1 + [\log_{10} k_0[M]/k_{\infty}]^2\right\}^{-1}$$

with

$$k_0 = 4 \times 10^{-30} \text{ cm}^6 \text{ molecule}^{-2} \text{ s}^{-1}$$

and

$$k_{m} = 8.7 \times 10^{-13} \text{ cm}^{3} \text{ molecule}^{-1} \text{ s}^{-1}$$

For the other alkynes studied to date, the rate constants determined at 760 torr total pressure of air are probably the high pressure limiting values.

Table 23. Room Temperature Rate Constants k for the Reaction of OH Radicals with Alkynes at the High-Pressure Limit (from Atkinson, 1986a)

Alkyne	$10^{12} \text{ x k (cm}^3 \text{ molecule}^{-1} \text{ s}^{-1})$
Acetylene	0.87
Propyne	5.8
1-Butyne	8.0
2-Butyne	29

These reactions proceed by initial OH radical addition to the $-C \equiv C - b$ bond. The OH-acetylene adduct can isomerize to the vinoxy (CH₂CHO) radical with subsequent decomposition or reaction. At low pressures, the formation of ketene has been observed, and this process may explain the observation of a bimolecular reaction with room temperature rate constant of $\sim (5 \pm 3) \times 10^{-14} \text{ cm}^3 \text{ molecule}^{-1}$ (Schmidt et al., 1985). Thus, the reaction probably proceeds via

For atmospheric purposes any "direct" formation of ketene is of negligible importance [with $\le 0.5\%$ of ketene being observed from acetylene (Hatakeyama et al., 1986)], with the majority of the reaction proceeding through the formation of a stabilized OH-acetylene adduct. Schmidt et al. (1985) observed the intermediate formation of the vinoxy radical, with glyoxal being a major product. In the presence of O_2 , OH radicals were efficiently regenerated (Schmidt et al., 1985),

$$CH_2CHO + O_2 + (CHO)_2 + OH$$

Hatakeyama et al. (1986) investigated the products of the reactions of OH and OD radicals with acetylene, propyne and 2-butyne. Glyoxal,

methylglyoxal and biacetyl, respectively, were observed products from these reactions in both the absence and presence of NO_X . For acetylene, formic acid was also observed in appreciable (40 \pm 10%) yield. The reaction of OH radicals with acetylene thus appears to proceed via (at the high pressure limit),

OH + CH=CH
$$\stackrel{*}{\leftarrow}$$
 [HOCH=CH]

[HOCH=CH] + M + HOCH=CH + M

HOCH=CH + CH₂CHO $\stackrel{\circ}{\rightarrow}$ (CHO)₂ + OH

HOCH=CH + O₂ + HOCH=CHOO

HOCH=CHOO + HCO + HCOOH

HOCH=CHOO + NO + HOCHCHO + NO₂
 \downarrow O₂

(CHO)₂ + HO₂

The amount of reaction leading directly to OH formation, without conversion of NO to NO_2 , is uncertain at the present time. The product yields obtained by Hatakeyama et al. (1986) were as follows: from acetylene, HCOOH, 0.4 \pm 0.1; (CHO $_2$), 0.7 \pm 0.3; from propyne, HCOOH, 0.12 \pm 0.02; CH $_3$ COCHO, 0.53 \pm 0.03; and from 2-butyne, CH $_3$ COOH, 0.12 \pm 0.01 and CH $_3$ COCOCH $_3$, 0.87 \pm 0.07. Other products must thus be formed from propyne.

03 Reactions

Kinetic data have been determined for the reactions of 0_3 with acetylene, propyne, 1-butyne, 2-butyne and butadiyne (Atkinson and Carter, 1984). There are significant discrepancies between the various studies, and no recommendations were made by Atkinson and Carter (1984). The most recent data of Atkinson and Aschmann (1984) give rate constants at 294 ± 2 K of (in cm³ molecule⁻¹ s⁻¹ units): acetylene, 7.8×10^{-21} ; propyne, 1.4 $\times 10^{-20}$, and 1-butyne, 2.0×10^{-20} . These rate constants are sufficiently low that the 0_3 reactions are of negligible importance as an alkyne loss process. No definitive product data are available, although α -dicarbonyls have been observed from the reaction of 0_3 with acetylene, propyne, 1-butyne and 2-butyne (DeMore, 1971).

NO3 Radical Reactions

Rate constants, or upper limits to the rate constants, have been determined at room temperature for the reaction of NO $_3$ radicals with acetylene and propyne (Atkinson et al., 1988b). Based upon a rate constant for the reaction of NO $_3$ radicals with ethene of 2.14 x 10 $^{-16}$ cm 3 molecule $^{-1}$ s $^{-1}$ (Atkinson et al., 1988b), rate constants (in cm 3 molecule $^{-1}$ s $^{-1}$ units) of ≤ 3.5 x 10 $^{-17}$ and 1.8 x 10 $^{-16}$ are obtained for acetylene and propyne, respectively, at 296 ± 2 K. These relative rate data are in reasonable agreement with the absolute rate constants at 295 ± 2 K of (5.1 ± 3.5) x 10 $^{-17}$ cm 3 molecule $^{-1}$ s $^{-1}$ and (2.66 ± 0.32) x 10 $^{-16}$ cm 3 molecule $^{-1}$ s $^{-1}$ for acetylene and propyne, respectively, reported by Canosa-Mas et al. (1986). These reactions are sufficiently slow that they can be neglected for atmospheric purposes.

VIII. OXYGEN-CONTAINING ORGANIC COMPOUNDS

In this section, the atmospheric chemistry of those oxygen-containing organic compounds which are either emitted into the troposphere directly or are formed in the atmosphere as degradation products of other organics is dealt with. The atmospheric chemistry of the simple aliphatic aldehydes, ketones and α -dicarbonyls has been discussed above in Section II.

A. Alcohols

The alcohols of interest in urban atmospheres are primarily methanol, ethanol and, to a lesser extent, the C_3 and C_4 species. The gas-phase reactions with the NO $_3$ radical are slow (Wallington et al., 1987a), with upper limits to the room temperature rate constants of $<6\times10^{-16}$, $<9\times10^{-16}$ and $<2.3\times10^{-15}$ cm 3 molecule $^{-1}$ s $^{-1}$ for methanol, ethanol and 2-propanol, respectively. While no kinetic data are available for the gas-phase reactions of O_3 with the alcohols (Atkinson and Carter, 1984), the reactions are expected to be of negligible importance as an atmospheric loss process. Thus the only loss process which requires consideration is that by reaction with the OH radical. The room temperature rate constants and temperature dependencies for the OH radical reactions with the alcohols are given in Table 24.

These OH radical reactions proceed by H-atom abstraction from both the C-H and O-H bonds. For methanol, the rate constant ratio $k_a/(k_a+k_b)$, where k_a and k_b are the rate constants for the reactions,

Table 24. Room Temperature Rate Constants k and Arrhenius Parameters ($k = Ae^{-E/RT}$) for the Gas-Phase Reactions of the OH Radical with a Series of Alcohols, Ethers, Carboxylic Acids, Esters, Epoxides, Hydroperoxides, α,β -Unsaturated Carbonyls and 1,4-Unsaturated Dicarbonyls

Organic	$10^{12} \times k \text{ (cm}^3$ molecule ⁻¹ s ⁻¹)	$10^{12} \times A \text{ (cm}^3$ molecule ⁻¹ s ⁻¹)	E/R(K)
Methanol ^a	0.90	9.1	690
Ethanol ^a	3.4	9.3	300
1-Propanol ^a	5.3		
2-Propanol ^a	5.6	5.6	0
1-Butanol ^b	8.3		
Dimethyl ether ^C	3.0	10.4	372
Diethyl ether ^C	13		
Methyl t-butyl ether ^d	3.1	5.1	155
Formic acid ^a	0.48	0.48	0
Acetic acid ^a	0.74	1.3	170
Methyl formate ^e	0.23		
Methyl acetate ^e	0.34	0.83	260
Ethyl acetate ^e	1.5	2.3	131
n-Propyl acetate ^e	3.5		
Ethene oxide ^C	0.081	11	1460
Propene oxide ^C	0.52		
Methyl hydroperoxide ^C	11		
t-Butyl hydroperoxide ^C	3.0		

(continued)

Table 24 (continued) - 2

Organic		$10^{12} \times A \text{ (cm}^3$ molecule ⁻¹ s ⁻¹)	E/R(K)
Acrolein ^C	20		
Methacrolein ^c	31	20	-134
Crotonaldehyde ^C	36		
Methyl vinyl ketone ^c	19	3.3	-514
cis-3-Hexene-2,5-dione	63		
trans-3-Hexene-2,5-dione	53		

aAtkinson et al. (1988a). bWallington and Kurylo (1987b). cAtkinson (1986a). dWallington et al. (1988a). eWallington et al. (1988b).

$$OH + CH_3OH \rightarrow H_2O + CH_3O$$
 (a)

$$OH + CH_3OH \rightarrow H_2O + CH_2OH$$
 (b)

have been determined to be 0.10-0.15 at 298 K (Atkinson, 1986a and references therein). For ethanol, Meier et al. (1985) have shown that the initial OH radical reaction forms CH₃CHOH 75 ± 15% of the overall reaction pathway at room temperature. The estimation technique of Atkinson (1986a,b) allows the distribution of initially formed radicals to be approximately determined for the higher alcohols, and the subsequent reactions of these radicals are then as discussed in Section I above.

B. Ethers

The ethers of main interest are dimethyl ether, diethyl ether, and methyl $\underline{\text{tert}}$ -butyl ether. The reactions of the aliphatic saturated ethers with 0_3 and 0_3 radicals are expected to be of negligible importance as atmospheric loss processes [with an upper limit to the rate constant for the reaction of 0_3 radicals with 0_3 of 0_3 of 0_3 x 0_3 molecule 0_3 molecule 0_3 at 298 K being the only available data (Wallington et al., 1986a)]. Hence the major, if not sole, atmospheric loss process will be by reaction with the OH radical. The available kinetic data for these reactions are given in Table 24. The expected reaction schemes (based upon the calculated values of 0_3 moleculated and methyl 0_3 moleculated in the presence of 0_3 moleculated values of 0_3 moleculated in the presence of 0_3 moleculated 0_3 moleculated values of 0_3 moleculated 0_3 moleculated

<u>Dimethyl ether:</u>

Methyl t-butyl ether

OH +
$$CH_3OC(CH_3)_3$$
 + H_2O + $CH_2OC(CH_3)_3$
+ H_2O + $CH_3OC(CH_3)_2CH_2$

followed by:

$$(CH_3)_3COCH_2 \xrightarrow{O_2, NO} (CH_3)_3COCH_2O$$
 $\downarrow O_2$
 $\downarrow O_2$
 $(CH_3)_3COCHO + HO_2$

$$CH_3OC(CH_3)_2CHO + HO_2$$

C. Carboxylic Acids

The carboxylic acids of interest in polluted urban atmospheres are formic and acetic acid. Again, although no data have been reported for the NO $_3$ radical or O $_3$ reactions, the only important gas-phase atmospheric removal process is expected to be by OH radical reaction. The OH radical rate constants are given in Table 24. For HCOOH, the OH radical reaction proceeds to form mainly CO $_2$ and an H-atom, although the reaction dynamics are not yet known. No product data are available for the OH radical reaction with CH $_3$ COOH. Since these OH radical reactions are slow, with calculated atmospheric lifetimes due to OH radical reaction of approximately 50 days, the major process removing these compounds from the gas phase will be by wet or dry deposition, with incorporation into raindrops or cloud and fog water being of importance from the viewpoint of acid deposition.

D. Esters

Analogous to the alcohols, ethers and carboxylic acids, the only important gas-phase atmospheric loss process for the simple esters is expected to be by OH radical reaction, and rate constants are given in Table 24 for the esters for which data are available. These data show that the reaction proceeds by H atom abstraction from the -OR group, for

example

$$OH + CH_3C(O)OCH_3 + H_2O + CH_3C(O)OCH_2$$

No product data for this class of organic compounds are available, but the subsequent reactions of the initially formed radicals are expected to be as discussed in Section II. For example, for the radical formed from methyl acetate, in the presence of NO:

$$CH_3C(0)OCH_2 + O_2 + CH_3C(0)OCH_2OO$$
.

 $NO \longrightarrow NO_2$
 $CH_3C(0)OCH_2O \xrightarrow{O_2} CH_3C(0)OCHO + HO_2$
 $HCHO + CH_3CO_2$
 $CH_3 + CO_2$

with the relative importance of $\mathbf{0}_2$ reaction versus decomposition of the alkoxy radical being uncertain at the present time.

E. Epoxides

The major epoxides of interest are ethene oxide and propene oxide.

Again, OH radical reaction is expected to be the dominant atmospheric removal process, and the rate constants for these reactions are given in Table 24 for those epoxides for which data are available. This OH radical

reaction will proceed by initial H atom abstraction

$$OH + CH_2 CH_2 + CH_2 CH_2 + H_2O$$

OH +
$$CH_3CH_2$$
 + H_2O + (mainly) CH_2CH_2

Only for ethene oxide has a product study been carried out. For the higher epoxides, the distribution of initially formed radicals can be estimated using the technique of Atkinson (1986b) with, as shown, propene oxide being calculated to yield mainly the 'CH₂CHCH₂ radical. Presumably due to the large amount of ring strain, the initially formed radical (if the radical center is a part of the 3-membered ring system) may rapidly undergo ring cleavage, for example:

$$cH_2$$
 + $[cH_2cHo]^{\ddagger}$ \xrightarrow{M} cH_2cHo

Lorenz and Zellner (1984) have observed the vinoxy (CH₂CHO) radical in appreciable (0.23 at 298 K and 60 torr total pressure of helium) yield from the OH radical reaction with ethene oxide. No product data are available for the higher epoxides.

F. Hydroperoxides

To date, data concerning the atmospheric chemistry of the hydroperoxides are only available for methyl hydroperoxide ($\mathrm{CH_300H}$) and $\mathrm{\underline{tert}}$ -butyl hydroperoxide [($\mathrm{CH_3}$)300H]. The expected gas-phase atmospheric loss processes for this class of organic compounds are by photolysis and OH radical reaction. The absorption cross section of $\mathrm{CH_300H}$ has been evaluated by NASA (DeMore et al., 1987), and the quantum yield is expected to be unity.

$$CH_3OOH + h\nu \rightarrow CH_3O + OH$$

The available room temperature rate constants for the OH radical reactions of CH_3OOH and $(\text{CH}_3)_3\text{COOH}$ are given in Table 24. For CH_3OOH , the study of Niki et al. (1983b) shows that the OH radical reaction proceeds by two pathways

$$OH + CH_3OOH \rightarrow H_2O + CH_2OOH$$
 (a)

$$OH + CH_3OOH + H_2O + CH_3OO$$
 (b)

with $k_a/(k_a+k_b)=0.4$ at 298 K. The CH₂OOH radical is expected to rapidly decompose

to regenerate the OH radical directly.

G. α, β-Unsaturated Carbonyls

The α ,8-Unsaturated carbonyls of major interest are acrolein, methacrolein and methyl vinyl ketone, these being formed from the atmospheric degradation reactions of 1,3-butadiene (acrolein) and 2-methyl-1,3-butadiene [isoprene] (methacrolein and methyl vinyl ketone). The major atmospheric removal processes for this class of organic compounds are expected to be reaction with 0_3 and with OH and 0_3 radicals. As shown by the recent data for acrolein (Gardner et al., 1987), photolysis appears to be of minor importance as a loss process [with a lifetime of acrolein of ~10 days at 0_3 zenith angle (Gardner et al., 1987)].

The kinetics of the gas-phase reactions of the NO $_3$ radical with acrolein and crotonaldehyde have been determined (Atkinson et al., 1988b), and the rate constants at 296 \pm 2 K are 1.2 x 10 $^{-15}$ and 5.1 x 10 $^{-15}$ cm 3 molecule $^{-1}$ s $^{-1}$, respectively. These reactions are expected to be analogous to the OH radical reactions (see below) in that the NO $_3$ radical reaction with acrolein is expected to proceed mainly by H-atom abstraction from the -CHO group:

$$NO_3 + CH_2 = CHCHO + HNO_3 + CH_2 = CHCO$$

while the reaction with crotonal dehyde will proceed by H-atom abstraction from the -CHO group and $\rm NO_3$ radical addition to the >C=C< bond

→ CH₃CH(ONO₂)CHCHO and

CH3CHCH(ONO2)CHO

These NO $_3$ radical reactions will be of generally minor significance as atmospheric loss processes of the α,β -unsaturated carbonyls. The room temperature rate constants for the gas-phase reactions of O_3 with acrolein, methacrolein, crotonaldehyde and methyl vinyl ketone have been measured (Atkinson and Carter, 1984) with rate constants of (in units of 10^{-18} cm 3 molecule $^{-1}$ s $^{-1}$) 0.28, 1.1, 0.9 and 4.8, respectively. As for the alkenes and haloalkenes, these O_3 reactions proceed by initial addition to the >C=C< bonds. Again, these O_3 reactions are of minor importance as atmospheric loss processes.

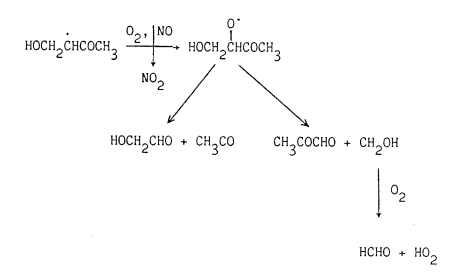
The major atmospheric loss process of the α,β -unsaturated carbonyls is with the OH radical, and the room temperature rate constants and Arrhenius parameters for this reaction for acrolein, methacrolein, crotonaldehyde and methyl vinyl ketone are given in Table 24. For the aldehydes, these OH radical reactions occur by H-atom abstraction from the -CHO group and OH radical addition to the >C=C< bond:

$$OH + CH_2 = CHCHO \rightarrow H_2O + CH_2 = CHCO$$
 (a)

For acrolein the major reaction pathway appears to be H-atom abstraction from the -CHO group, while for crotonaldehyde and methacrolein both reaction pathways occur (Atkinson, 1986a). The subsequent reactions of the initially formed radicals will be as discussed in Section I.

For the α , β -unsaturated ketones such as methyl vinyl ketone the only reaction pathway (since H-atom abstraction from the substituent alkyl groups will be slow) involves OH radical addition to the >C=C< bond:

The subsequent reactions of these radicals are as dealt with in Section I; i.e., in the presence of NO



Formation of $HOCH_2CHO + CH_3CO$ is calculated (Benson, 1976) to be ~10 kcal mol^{-1} more exothermic than the formation of $CH_3COCHO + CH_2OH$, and this is in accord with the product data of Tuazon et al. (unpublished data, 1988).

H. 1,4-Unsaturated Carbonyls

The 1,4-unsaturated carbonyl compounds have been postulated to be formed in the atmospheric degradation of the aromatic hydrocarbons (see, for example, Atkinson and Lloyd, 1984), and 3-hexene-2,5-dione $(CH_3COCH=CHCOCH_3)$ has been observed in small yield from the photooxidation of 1,2,4-trimethylbenzene (Takagi et al., 1982) and, in larger yield, from p-xylene (Becker and Klein, 1987). Experimental data are only available for the atmospherically important reactions of cis- and trans-3-hexene-2,5-dione (Tuazon et al., 1985; Becker and Klein, 1987). These isomers were observed to react with 0_3 , with rate constants at 298 ± 2 K of 1.8 x 10^{-18} and 8.3×10^{-18} cm³ molecule⁻¹ s⁻¹ for <u>cis</u>- and <u>trans</u>-3-hexene-2,5dione, respectively (Tuazon et al., 1985). The measured OH radical reaction rate constants are given in Table 24. Photolysis of both isomers was also observed, with the major pathway in the spectral region employed (>320 nm) being photoisomerization to the other isomer (Tuazon et al., 1985; Becker and Klein, 1987). Based upon these data, it appears that photolysis (other than photoisomerization) and reaction with the OH radical will be the dominant atmospheric removal processes.

Unfortunately, no definitive product data have been obtained for the OH radical reactions with these 1,4-unsaturated carbonyls, although a speculative discussion of the likely reactions of this class of organic compounds and their rates and mechanisms has been given by Atkinson and Lloyd (1984). However, until it can be determined whether or not these compounds are indeed formed to any significant extent from the degradation of aromatic hydrocarbons or other organic emissions, speculation about the atmospheric reactions of these chemicals is not warranted.

IX. SULFUR-CONTAINING ORGANIC COMPOUNDS

The organosulfur compounds which need to be considered are methyl mercaptan (CH₃SH), dimethyl sulfide (CH₃SCH₃) and dimethyl disulfide (CH₃SSCH₃). The atmospherically important reactions of these organosulfur species have been evaluated by the IUPAC data panel (Atkinson et al., 1988a), and the recommendations of that evaluation are used here. The potentially significant atmospheric loss processes for these chemicals are by reaction with OH and NO₃ radicals and with O₃. Photolysis (Calvert and Pitts, 1966) is expected to be of negligible importance due to the high reactivity of these compounds toward the OH radical. Only for dimethyl sulfide has the kinetics of the O₃ reaction been studied, with an upper limit for this reaction of 8.3 x 10^{-19} cm³ molecule⁻¹ s⁻¹ at room temperature (Atkinson and Carter, 1984). Atkinson and Carter (1984) recommend that the gas-phase reactions of O₃ with these organosulfur compounds be considered to be unimportant as an atmospheric loss process, and thus only the OH and NO₃ radical reactions need to be dealt with.

OH Radical Reactions

The room temperature rate constants and Arrhenius parameters for the OH radical reactions with $\mathrm{CH_3SH}$, $\mathrm{CH_3SCH_3}$ and $\mathrm{CH_3SSCH_3}$ are given in Table 25. For $\mathrm{CH_3SH}$ (and $\mathrm{CD_3SH}$), no effect of either total pressure or of the presence of $\mathrm{O_2}$ on the rate constant has been observed (Hynes and Wine, 1987), and a simple Arrhenius expression is sufficient. The lack of a kinetic isotope effect (Atkinson, 1986a; Hynes and Wine, 1987) and the product data of Hatakeyama and Akimoto (1983) show that this OH radical reaction with $\mathrm{CH_3SH}$ proceeds by initial addition of the OH radical to the S atom, followed by reactions to yield $\mathrm{SO_2}$, $\mathrm{CH_3SO_3H}$ (methanesulfonic acid)

Table 25. Room Temperature Rate Constants k and Arrhenius Parameters (k = $Ae^{-E/RT}$) for the Gas-Phase Reactions of the OH Radical with Selected Organosulfur Compounds (from Atkinson et al., 1988a, except as indicated)

Organic	10^{12} x k $(\text{cm}^3 \text{ molecule}^{-1} \text{ s}^{-1})$	$10^{12} \times A$ (cm ³ molecule ⁻¹ s ⁻¹)	E/R(K)
CH ₃ SH	33	9.9	-356
сн ₃ sсн ₃	$4.4 + \frac{(4.1 \times 10^{-17}[0_2])}{(1 + 4.1 \times 10^{-20}[0_2])}$	a	· a
сн ₃ sscн ₃ b	205	51	-414

^aSee text. ^bFrom Atkinson (1986a).

and HCHO. Hatakeyama and Akimoto (1983) suggested that the initial adduct decomposes to CH_3S and H_2O , followed by reactions of CH_3S radicals.

OH +
$$CH_3SH \rightarrow CH_3S(OH)H$$

$$CH_3S + H_2O$$

In the presence of alkyl nitrites (used to generate OH radicals in the experimental system used), Hatakeyama and Akimoto (1983) observed the formation of CH_3SNO , with the N atom coming from the alkyl nitrite (and not from NO), with subsequent photolysis of CH_3SNO giving rise to CH_3S radicals and hence to the observed products. The formation of CH_3SNO was attributed to the reaction of the $\text{CH}_3\text{S}(\text{OH})\text{H}$ adduct with the alkyl nitrite

$$CH_3S(OH)H + RONO \rightarrow CH_3SNO + RCH_2OH + OH$$

For $\mathrm{CH_3SSCH_3}$, the OH radical reaction also appears to proceed by OH radical addition (Hatakeyama and Akimoto, 1983; Atkinson, 1986a), leading to the production of $\mathrm{CH_3S}$ and $\mathrm{CH_3SOH}$ radicals, the subsequent reactions of which lead to the $\mathrm{SO_2}$, HCHO and $\mathrm{CH_3SO_3H}$ products.

OH +
$$CH_3SSCH_3$$
 + $[CH_3SS(OH)CH_3]$

$$CH_3S + CH_3SOH$$

In contrast, for CH_3SCH_3 the OH radical reaction proceeds by two pathways; H atom abstraction from the C-H bonds and OH radical addition to the S atom

$$OH + CH_3SCH_3 \rightarrow H_2O + CH_3SCH_2$$
 (a)

$$\rightarrow \text{CH}_3\text{S}(\text{OH})\text{CH}_3$$
 (b)

The recent kinetic data of Hynes et al. (1986) [which were, in a preliminary form, incorporated and discussed in the review of Atkinson (1986a)] show that in the absence of O_2 the OH-CH₃SCH₃ adduct rapidly back-decomposes to the reactants. Thus, in the absence of O_2 the only reaction pathway observed is the H atom abstraction route, with a rate constant of (Atkinson et al., 1988a)

$$k_a = 9.6 \times 10^{-12} e^{-234/T} cm^3 molecule^{-1} s^{-1}$$

$$= 4.4 \times 10^{-12} \text{ cm}^3 \text{ molecule}^{-1} \text{ s}^{-1} \text{ at } 298 \text{ K}.$$

In the presence of $\mathrm{O_2}$, the $\mathrm{OH-CH_3SCH_3}$ adduct reacts

$$CH_3S(OH)CH_3 + O_2 \rightarrow products$$

Thus, the observed rate constant for the OH radical addition pathway is dependent on the $\rm O_2$ concentration, and based upon the kinetic data of Hynes et al. (1986), Atkinson et al. (1988a) recommend that

$$k_b = \frac{1.7 \times 10^{-42} e^{7810/T} [O_2]}{(1 + 5.5 \times 10^{-31} e^{7460/T} [O_2])} cm^3 \text{ molecule}^{-1} s^{-1}$$

over the temperature range ~260-360 K. Hence, at 298 K and atmospheric pressure (760 torr) of air,

$$k(OH + CH_3SCH_3) = k_a + k_b = 6.2 \times 10^{-12} \text{ cm}^3 \text{ molecule}^{-1} \text{ s}^{-1}$$

The H-atom abstraction pathway is expected to proceed by the reactions

OH +
$$CH_3SCH_3 \rightarrow H_2O + CH_3SCH_2$$

NO $\downarrow O_2$
 CH_3SCH_2O
 $\downarrow CH_3SCH_2O$
 $\downarrow CH_3SCH_2O$

followed by reactions of the CH_3S radical (discussed below). However, the reactions subsequent to the formation of the $CH_3S(OH)CH_3$ radical, in the presence of O_2 , are not known at present. Furthermore, the complexities of the experimental systems used to investigate the products of the OH_2SCH_3 reaction under simulated atmospheric conditions [involving secondary reactions leading to enhanced loss processes of CH_3SCH_3 not involving OH radicals (Atkinson, 1986a; Wallington et al., 1986b)] do not allow the products of the OH radical reactions to be quantitatively determined.

NO3 Radical Reactions

The kinetics of the gas-phase reactions of the NO₃ radical with several organosulfur species have recently been measured, and the rate constants recommended by Atkinson et al. (1988a) are given in Table 26 for CH₃SH, CH₃SCH₃ and CH₃SSCH₃. No definitive product or mechanistic data are available, but is appears that these reactions proceed by initial NO₃ radical addition, probably followed by rapid decomposition to, among other products, CH₃S radicals (Mac Leod et al., 1986). Speculative, and incomplete, reaction sequences have been presented by Mac Leod et al. (1986), and that reference should be consulted (although it must be recognized that much more work is necessary before any realistic reaction schemes can be proposed for use in urban airshed chemical mechanisms).

Reactions of the CH3S Radical

The ${\rm CH_3S}$ radical is becoming recognized as a key intermediate in the atmospheric degradation reactions of organosulfur compounds, leading to the formation of ${\rm SO_2}$, ${\rm CH_3SO_3H}$ and HCHO, and the atmospheric reactions of ${\rm CH_3S}$ thus need to be understood. To date, rate constants for the reactions of the ${\rm CH_3S}$ radical with NO, ${\rm NO_2}$, ${\rm O_2}$ and ${\rm O_3}$ have been reported (Balla et al., 1986; Black and Jusinski, 1986). The reaction with NO is an addition reaction

and the rate constant is in the fall-off regime between second- and third-order kinetics below atmospheric pressure for M = N_2 (Balla et al.,

Table 26. Room Temperature Rate Constants k and Arrhenius Parameters $(k = Ae^{-E/RT})$ for the Gas-Phase Reactions of the NO $_3$ Radical with Selected Organosulfur Compounds (from Atkinson et al., 1988a)

Organic	10^{12} x k $(\text{cm}^3 \text{ molecule}^{-1} \text{ s}^{-1})$	$10^{12} \times A$ (cm ³ molecule ⁻¹ s ⁻¹)	E/R(K)
СН ₃ SH	1.0	1.0	0
CH ₃ SCH ₃	1.0	0.19	- 500
сн ₃ sscн ₃	0.7	0.7	0

1986). At 295 K, Balla et al. (1986) fit their data for M = $\rm N_2$ to the Troe expression

$$k = \left(\frac{k_o[M]}{1 + \frac{k_o[M]}{k_{\infty}}}\right) F \left\{1 + \left[\log \left(k_o[M]/k_{\infty}\right)\right]^2\right\}^{-1}$$

with

$$k_o = 3.3 \times 10^{-29} \text{ cm}^6 \text{ molecule}^{-2} \text{ s}^{-1},$$

 $k_{\infty} = 3.6 \times 10^{-11} \text{ cm}^3 \text{ molecule}^{-1} \text{ s}^{-1}$

and

$$F = 0.60.$$

At the high pressure limit (determined using SF_6 as the diluent gas),

$$k(CH_3S + NO) = 1.81 \times 10^{-12} e^{906/T} cm^3 molecule^{-1} s^{-1}$$

The reaction of CH_3S radicals with NO_2 exhibits bimolecular kinetics with no marked pressure dependence, and the rate constant determined by Balla et al. (1986) is

$$k(CH_3S + NO_2) = 8.3 \times 10^{-11} e^{81/T} cm^3 molecule^{-1} s^{-1}$$

$$= 1.1 \times 10^{-10} \text{ cm}^3 \text{ molecule}^{-1} \text{ s}^{-1} \text{ at } 298 \text{ K}.$$

It is possible that this reaction forms $\text{CH}_3\text{SO} + \text{NO}$. The reaction of CH_3S radicals with O_2 is slow, with an upper limit to the rate constant of <2 x 10^{-17} cm³ molecule⁻¹ s⁻¹ at room temperature (Balla et al., 1986). Black

and Jusinski (1986) have reported, using a somewhat complex chemical system, an upper limit to the reaction of CH $_3$ S radicals with O $_3$ of <8 x $_{10^{-14}~\rm cm}^3$ molecule $^{-1}~\rm s^{-1}$ at room temperature.

Although further kinetic and product data are clearly required for these reactions of the ${\rm CH_3S}$ radical, it appears that the major reaction in the polluted atmosphere will be with ${\rm NO_2}$ [since the ${\rm CH_3SNO}$ formed from the NO reaction will rapidly photodissociate back to ${\rm CH_3S}$ + NO (Niki et al., 1983c) during daylight]. The situation is less clear for the "clean" troposphere until further, more definitive, rate data become available for the reaction of ${\rm CH_3S}$ radicals with ${\rm O_2}$.

X. NITROGEN-CONTAINING ORGANIC COMPOUNDS

The nitrogen-containing organic compounds considered in this section are the simple aliphatic amines (and related alcohol amines), the alkyl nitrates expected to be formed in the atmosphere from the alkanes (see Section III), and peroxyacyl and peroxyalkyl nitrates. These are dealt with below.

Aliphatic Amines

The expected atmospheric loss processes for these compounds are by gas-phase reaction with nitric acid (HNO $_3$), OH and NO $_3$ radicals and O $_3$. The amines will react with gaseous HNO $_3$, in a manner similar to NH $_3$, to form their nitrate salts

$$RNH_2 + HNO_3 \rightarrow RNH_3^+ NO_3^-$$

although no rate data are currently available. Similarly, although it is expected that NO_3 radicals may react rapidly with the amines, no

experimental data are available due to difficulties in studying these reactions in the absence of \mbox{HNO}_3 .

The gas-phase reactions of the simple aliphatic amines with 0_3 have been studied (Atkinson and Carter, 1984), and the measured room temperature rate constants range from 2 x 10^{-20} cm³ molecule⁻¹ s⁻¹ for methylamine to 1.0 x 10^{-17} cm³ molecule⁻¹ s⁻¹ for trimethylamine (although these rate constants must be viewed as upper limits because of the possibility if secondary reactions consuming the reactants). Due to the high rate constants for the OH radical reactions (see below), these 0_3 reactions are calculated to be of minor significance under atmospheric conditions.

The rate constants for the OH radical reactions with the amines and alcohol amines are given in Table 27. While the reaction products are those corresponding to H atom abstraction, these reactions with the amines probably occur by initial OH radical addition to the N atom, followed by rapid decomposition of this adduct to products. For the reaction of OH radicals with $(CH_3)_2NH$, Lindley et al. (1979) determined the rate constant ratio $k_a/(k_a+k_b)=0.37\pm0.05$ at room temperature, where k_a and k_b are the rate constants for the reaction pathways (a) and (b), respectively.

$$OH + (CH_3)_2NH + H_2O + (CH_3)_2\dot{N}$$
 (a)

$$OH + (CH_3)_2NH + H_2O + CH_3NH\dot{C}H_2$$
 (b)

From these data and the kinetics of these reactions, it appears that for methylamine and ethylamine the OH radical reactions proceed by H atom abstraction from the N-H bonds, for dimethylamine H atom abstraction

Table 27. Room Temperature Rate Constants k and Arrhenius Parameters $(k = Ae^{-E/RT})$ for the Gas-Phase Reactions of the OH Radical with Selected Nitrogen-Containing Compounds [from Atkinson (1986a) except as indicated]

Organic	$10^{12} \times k \text{ (cm}^3$ $\text{molecule}^{-1} \text{ s}^{-1}$)	$10^{12} \times A \text{ (cm}^3$ molecule ⁻¹ s ⁻¹)	E/R(K)
Methylamine	22	10	-229
Ethylamine	28	15	-189
Dimethylamine	65	29	-247
Triethylamine	61	26	-252
2-Dimethylaminoethanol	80	80	0
2-Amino-2-methyl-1-propanol	28		
N-Nitrosodimethylamine	2.5		
Dimethylnitramine	3.8		
Methyl nitrate ^a	0.034		
Ethyl nitrate ^b	0.49		•
1-Propyl nitrate ^b	0.49		
2-Propyl nitrate	0.18		
1-Butyl nitrate	1.4		
2-Butyl nitrate	0.67		
2-Pentyl nitrate	1.8		
3-Pentyl nitrate	1.1		
2-Methyl-3-butyl nitrate	1.7		
2,2-Dimethyl-1-propyl nitrate	0.85		
2-Hexyl nitrate	3.1		

(continued)

Table 27 (continued) - 2

Organic		$10^{12} \times A \text{ (cm}^3$ molecule ⁻¹ s ⁻¹)	E/R(K)
3-Hexyl nitrate	2.7		
Cyclohexyl nitrate	3.3		
2-Methyl-2-pentyl nitrate	1.7		
3-Methyl-2-pentyl nitrate	3.0		
3-Heptyl nitrate	3.6		
3-Octyl nitrate	3.8		
Peroxyacetyl nitrate	0.14	1.2	650

^aGaffney et al. (1986). ^bKerr and Stocker (1986).

occurs from both the N-H and C-H bonds, and for trimethylamine the reaction proceeds by H atom abstraction from the C-H bonds. Atkinson (1986b) has formulated an estimation technique for the calculation of OH radical reaction rate constants (and the positions at which reaction occurs) for these related compounds.

The subsequent reactions are not totally understood at present, but Lindley et al. (1979) showed that for the reactions of the dimethylamino radical with 0_2 , NO and $N0_2$,

$$(CH_3)_2N + O_2 \xrightarrow{M} (CH_3)_2NOO$$

 $+ CH_3N=CH_2 + HO_2$ (a)

$$(CH_3)_2N + NO + (CH_3)_2NNO$$
 (b)
 $(N-nitrosodimethylamine)$

$$(CH_3)_2^{N} + NO_2^{+} + (CH_3)_2^{NNO_2}$$
 (c)
(dimethylnitramine)

$$+ CH_3N=CH_2 + HONO$$
 (d)

 $k_{\rm d}/k_{\rm c}$ = 0.22, $k_{\rm a}/k_{\rm c}$ = 3.9 x 10⁻⁷ and $k_{\rm a}/k_{\rm b}$ = 1.5 x 10⁻⁷ at room temperature.

Thus, under polluted urban atmospheric conditions the major reaction of the dimethylamino radical will be with NO and/or NO $_2$. For the CH $_3$ NHCH $_2$ radical also formed from dimethylamine, Lindley et al. (1979) showed that reaction with O $_2$ occurs via

$$CH_3NHCH_2 + O_2 + HO_2 + CH_3N=CH_2$$

The atmospheric reactions of N-nitrosodimethylamine and dimethylnitramine have been studied by Tuazon et al. (1983). The reactions of 0_3 with these compounds are slow, with measured upper limits of $<1 \times 10^{-20} \text{ cm}^3$ molecule⁻¹ s⁻¹ for N-nitrosodimethylamine and $<3 \times 10^{-21} \text{ cm}^3$ molecule⁻¹ s⁻¹ for dimethylnitramine at 296 K. The room temperature rate constants for the OH radical reactions are given in Table 27, with the reactions being expected to proceed by H atom abstraction from the C-H bonds. For N-nitrosodimethylamine photolysis also occurs

$$(CH_3)_2NNO + hv \rightarrow (CH_3)_2N + NO$$

with a lifetime of approximately 5 min at 0° zenith angle (Tuazon et al., 1983). Thus, the major atmospheric loss process for dimethylnitramine will be by OH radical reaction, while that for N-nitrosodimethylamine will be photolysis back to its precursors.

Alkyl Nitrates

Based upon laboratory studies concerning the formation of alkyl nitrates from the NO_{X} -air photooxidations of the alkanes, the major alkyl nitrates expected to be formed in the atmosphere are 2-propyl nitrate, 2-butyl nitrate, 2- and 3-pentyl nitrate, and 2- and 3-hexyl nitrate. Although no experimental data are available concerning the kinetics of the NO_3 radical or O_3 reactions, these are expected to be of no significance as atmospheric loss processes. The major loss processes are then reaction

with the CH radical and photolysis. The room temperature OH radical reaction rate constants are given in Table 27 (no temperature dependent data are available). These reactions proceed by H atom abstraction from the C-H bonds, and in the absence of product data the estimation technique of Atkinson (1987) can be used to calculate the distribution of the nitratoalkyl radicals formed in these reactions. For example

$$^{\circ}_{+}$$
 H₂0 + CH₃CHCH₂CHCH₃ (61%)

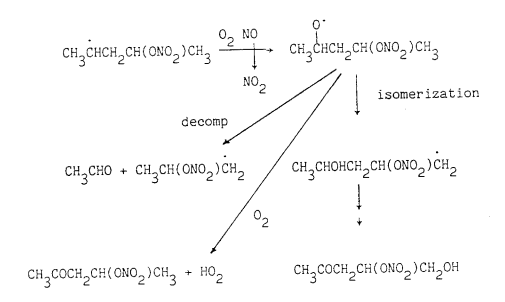
$$^{\text{ONO}_2}$$
 + $^{\text{H}_2\text{O}}$ + $^{\text{CH}_3\text{CH}_2\text{CHCHCH}_3}$ (13%)

$$+ \text{H}_2\text{O} + \text{CH}_3\text{CH}_2\text{CH}_2\text{C(ONO)}_2\text{CH}_3$$
 (13%)

$$^{ONO}_{1}^{2}$$

+ $^{H}_{2}^{O}$ + $^{CH}_{3}^{CH}_{2}^{CH}_{2}^{CH}_{2}^{CH}_{2}$ (2%)

The expected subsequent reactions of these radicals are, in the presence of $\mathrm{NO}_{\mathrm{X}},$ for example



with isomerization being expected to dominate (Section I)

$$\begin{array}{c} \text{CH}_3\text{CH}_2\text{CHCH}(\text{ONO}_2)\text{CH}_3 & \begin{array}{c} \text{O}_2 & \text{NO} \\ \text{NO}_2 & \end{array} \\ & \begin{array}{c} \text{CH}_3\text{CH}_2\text{CHCHCH}_3 \\ \text{ONO}_2 & \end{array} \\ & \begin{array}{c} \text{CH}_3\text{CHO} + \text{CH}_3\text{CHONO}_2 \\ \text{fast} \\ \end{array} \\ & \begin{array}{c} \text{CH}_3\text{CHO} + \text{NO}_2 \\ \end{array}$$

The absorption cross sections of methyl nitrate have been measured by Taylor et al. (1980) and, assuming a quantum yield of unity (consistent with the broad featureless nature of the absorption band), an atmospheric lifetime of approximately 10 days due to photolysis is calculated. The absorption band positions and cross sections of the higher alkyl nitrates are probably similar (Calvert and Pitts, 1966), leading to similar photolysis lifetimes to that for methyl nitrate. Photolysis will lead to

cleavage of the $O-NO_2$ bond

$$RONO_2 + hv + RO + NO_2$$

followed by the reactions of the alkoxy radical (Section I).

Peroxyacyl Nitrates and Peroxy Nitrates

Peroxyacetyl nitrate (PAN; CH₃C(0)00NO₂) and the higher members of this class of compounds are formed from the atmospheric degradation of many organics (see Section I). The reaction of OH radicals with PAN (the only peroxyacyl nitrate studied to date) is slow at tropospheric temperatures (Table 27) and this OH radical reaction is expected to be a significant atmospheric loss process for PAN only at the low temperatures encountered in the upper troposphere (Wallington et al., 1984). The major, and usually totally dominant, loss process for PAN in the atmosphere is by its thermal decomposition

$$CH_3C(0)OONO_2 + CH_3C(0)OO + NO_2$$

Thus, PAN is in equilibrium with NO_2 and the $CH_3C(0)00$ radical. In the presence of NO, the reaction

$$CH_3C(0)00 + NO + CH_3C(0)0 + NO_2$$

$$CH_3 + CO_2$$

leads to loss of PAN. The thermal decomposition rate constants for PAN are temperature and pressure dependent (Reimer and Zabel, 1986), being in the fall-off regime between first- and second-order kinetics below atmospheric pressure at room temperature. Using the Troe expression,

$$k = \left(\frac{k_{o}[M]}{\frac{k_{o}[M]}{1 + \frac{k_{o}[M]}{k_{\infty}}}}\right) F \left\{1 + [N^{-1} \log k_{o}[M]/k_{\infty}]^{2}\right\}^{-1}$$

Reimer and Zabel (1986) derived (for $M=N_2$)

$$k_0 = 6.3 \times 10^{-2} e^{-12783/T} cm^3 molecule^{-1} s^{-1}$$

$$k_m = 2.2 \times 10^{16} e^{-13437/T} s^{-1}$$

$$F = 0.27$$

and

$$N = 1.47$$

with

$$k_0^{0} 2/k_0^{N} 2 = 0.9 \pm 0.2$$

At 298 K and 760 torr total pressure of air,

$$k(PAN) = 4.2 \times 10^{-4} \text{ s}^{-1}$$

in good agreement with the previous recommendation of Atkinson and Lloyd (1984) of 3.6 x 10^{-4} s⁻¹ and with the atmospheric pressure data of

Schurath and Wipprecht (1980) [k = $3.2 \times 10^{16} \ e^{-13592/T} \ s^{-1}$ at atmospheric pressure, = $5.0 \times 10^{-4} \ s^{-1}$ at 298 K and atmospheric pressure]. The decomposition rates of the higher peroxyacyl nitrates are expected to be similar to that for PAN, although of course the decompositions will be closer to the high pressure limit at a given pressure than that for PAN, and at atmospheric pressure Schurath and Wipprecht (1980) determined that for peroxypropionyl nitrate (PPN; $C_2H_5C(0)00NO_2$),

$$k(PPN) = 1.6 \times 10^{17} e^{-14073/T} s^{-1}$$

= $5.0 \times 10^{-4} \text{ s}^{-1}$ at 298 K and atmospheric pressure.

For the peroxyalkyl nitrates, ROONO2, the only significant atmospheric loss process is again thermal decomposition

$$ROONO_2 \rightarrow RO_2 + NO_2$$

with the thermal decomposition rate constant for CH_3OONO_2 being well into the fall-off regime at pressures of one atmosphere and below (Reimer and Zabel, 1986). Using the Troe fall-off expression

$$k = \left(\frac{k_0[M]}{\frac{k_0[M]}{1 + \frac{k_0[M]}{k_{\infty}}}}\right) F \left\{1 + [N^{-1} \log k_0[M]/k_{\infty}]^2\right\}^{-1}$$

Reimer and Zabel (1986) derived for M=N2

$$k_0 = 8.5 \times 10^{-4} e^{-10317/T} cm^3 molecule^{-1} s^{-1}$$

$$k_{\infty} = 2.3 \times 10^{16} e^{-10770/T} s^{-1}$$

$$F = 0.47$$

and

$$N = 0.75 - 1.27 \log F = 1.17$$

with

$$k_0^{0}2/k_0^{0}2 = 1.2 \pm 0.2$$

At 760 torr total pressure of air and 298 K,

$$k(CH_3OONO_2) = 2.2 s^{-1}$$

The decomposition rates for the higher $ROONO_2$ species are expected to be similar to that for CH_3OONO_2 , except that they will be closer to the high pressure first-order limit (which is 4.8 s⁻¹ for CH_3OONO_2 at 298 K).

XI. AROMATIC COMPOUNDS

The aromatic compounds of concern in polluted urban atmospheres are the aromatic hydrocarbons and their aromatic ring-retaining products (aromatic aldehydes and phenolic compounds). In addition, there is recent concern over the atmospheric chemistry of the gas-phase polycyclic aromatic hydrocarbons. These are dealt with in this section.

Aromatic Hydrocarbons

The aromatic hydrocarbons react only very slowly with 0_3 and with $N0_3$ radicals, with room temperature rate constants for the 0_3 reactions of <1

x 10⁻²⁰ cm³ molecule⁻¹ s⁻¹ (Atkinson and Carter, 1984) and for the NO₃ radical reactions of 10⁻¹⁷ to 10⁻¹⁶ cm³ molecule⁻¹ s⁻¹ (Atkinson et al., 1988b). Thus, atmospheric removal of the aromatic hydrocarbons by these reactions is of no importance. The only important atmospheric loss process is by reaction with the OH radical, and the room temperature rate constants for this reaction are given in Table 28. In general, the temperature dependencies of these rate constants are small below room temperature (Atkinson, 1986a). These OH radical reactions proceed by two routes; H atom abstraction from the C-H bonds of the alkyl substituent groups (or the ring C-H bonds for benzene) and OH radical addition to the aromatic ring

OH +
$$CH_3$$
 CH_2 (a)

$$CH_3$$

$$CH_3$$

$$OH$$

$$CH_3$$

$$OH$$

$$(b)$$

$$(plus other isomers)$$

The H atom abstraction route is of minor importance, and the measured or estimated rate constant ratios $k_a/(k_a+k_b)$ at room temperature are also given in Table 28 (these are probably uncertain to a factor of approximately 2 for benzene, the xylenes and the trimethylbenzenes).

The reactions subsequent to the H atom abstraction route from the alkyl side chains are expected to be consistent with the data given in Section I, for example for the benzyl radical formed from toluene in the presence of NO

Table 28. Room Temperature Rate Constants for the Gas-Phase Reactions of the OH Radical with Aromatic Compounds and Rate Constant Ratios $k_a/(k_a+k_b)$

Aromatic (cm3 molecule-1 s-1) k _a /(k _a + k _b) Benzene 1.3 ^a 0.07 ^a Toluene 6.0 ^a , ^d 0.11 ^a , 0.08 ^b Ethylbenzene 7 ^a Isopropylbenzene 7 ^a Isopropylbenzene 5 ^a o-Xylene 14a, ^d 0.09 ^a , 0.05 ^c m-Xylene 24a, ^d 0.04 ^a , 0.04 ^c p-Xylene 12 ^a m-Ethyltoluene 12 ^a m-Ethyltoluene 12 ^a 1,2,3-Trimethylbenzene 33 ^d 0.05 ^a 1,2,4-Trimethylbenzene 33 ^d 0.04 ^a 1,3,5-Trimethylbenzene 58 ^d 0.03 ^a Benzaldehyde 13 ^a Phenol 28 ^a o-Cresol 40 ^a 0.08 ^a p-Cresol 40 ^a 0.08 ^a		10 ¹² x k	
Toluene 6.0a,d 0.11a, 0.08b Ethylbenzene 7a n-Propylbenzene 7a Isopropylbenzene 7a t-Butylbenzene 5a 0-Xylene 14a,d 0.09a, 0.05c m-Xylene 24a,d 0.04a, 0.04c p-Xylene 14a,d 0.09a, 0.08c o-Ethyltoluene 12a m-Ethyltoluene 12a 1,2,3-Trimethylbenzene 33d 0.05a 1,2,4-Trimethylbenzene 33d 0.04a 1,3,5-Trimethylbenzene 58d 0.03a Benzaldehyde 13a Phenol 28a 0-Cresol 40a 0.08a	Aromatic	•	$k_a/(k_a + k_b)$
Ethylbenzene 7a n-Propylbenzene 7a Isopropylbenzene 7a t-Butylbenzene 5a o-Xylene 14a,d 0.09a, 0.05c m-Xylene 24a,d 0.04a, 0.04c p-Xylene 12a m-Ethyltoluene 12a m-Ethyltoluene 12a t,2,3-Trimethylbenzene 33d 0.05a 1,2,4-Trimethylbenzene 58d 0.03a Benzaldehyde 13a Phenol 28a o-Cresol 40a 0.08a m-Cresol 57a m-Bathylbenzene 58d m-Cresol 57a m-Cresol 57a	Benzene	1.3 ^a	
Table Tabl		6.0 ^a ,d	0.11 ^a , 0.08 ^b
n-Propylbenzene 7a Isopropylbenzene 7a t-Butylbenzene 5a o-Xylene 14a,d 0.09a, 0.05c m-Xylene 24a,d 0.04a, 0.04c p-Xylene 14a,d 0.09a, 0.08c o-Ethyltoluene 12a 0.09a, 0.08c p-Ethyltoluene 12a 0.05a 1,2,3-Trimethylbenzene 33d 0.05a 1,2,4-Trimethylbenzene 33d 0.04a 1,3,5-Trimethylbenzene 58d 0.03a Benzaldehyde 13a 0.08a Phenol 28a 0.08a o-Cresol 40a 0.08a m-Cresol 40a 0.08a	Ethylbenzene	7 ^a	
Isopropylbenzene 7a t-Butylbenzene 5a o-Xylene 14a,d 0.09a, 0.05c m-Xylene 24a,d 0.04a, 0.04c p-Xylene 14a,d 0.09a, 0.08c o-Ethyltoluene 12a 0.09a, 0.08c m-Ethyltoluene 12a 0.05a 1,2,3-Trimethylbenzene 33d 0.05a 1,2,4-Trimethylbenzene 33d 0.04a 1,3,5-Trimethylbenzene 58d 0.03a Benzaldehyde 13a 0.08a Phenol 28a 0.08a o-Cresol 40a 0.08a m-Cresol 57a 0.08a	•	7 ^a	
o-Xylene 14a,d 0.09a, 0.05c m-Xylene 24a,d 0.04a, 0.04c p-Xylene 14a,d 0.09a, 0.05c o-Ethylene 14a,d 0.09a, 0.08c o-Ethylene 12a m-Ethylene 20a p-Ethylene 12a 1,2,3-Trimethylbenzene 33d 0.05a 1,2,4-Trimethylbenzene 33d 0.04a 1,3,5-Trimethylbenzene 58d 0.03a Benzaldehyde 13a Phenol 28a o-Cresol 40a 0.08a m-Cresol 57a		7 ^a	
o-Xylene 14a,d 0.09a, 0.05c m-Xylene 24a,d 0.04a, 0.04c p-Xylene 14a,d 0.09a, 0.04c o-Ethyltoluene 12a m-Ethyltoluene 12a 1,2,3-Trimethylbenzene 33d 0.05a 1,2,4-Trimethylbenzene 33d 0.04a 1,3,5-Trimethylbenzene 58d 0.03a Benzaldehyde 13a Phenol 28a o-Cresol 40a 0.08a m-Cresol 57a	t-Butylbenzene	5 ^a	
m-Xylene 24a,d 0.04a, 0.04c p-Xylene 14a,d 0.09a, 0.08c o-Ethyltoluene 12a m-Ethyltoluene 12a 1,2,3-Trimethylbenzene 33d 0.05a 1,2,4-Trimethylbenzene 33d 0.04a 1,3,5-Trimethylbenzene 58d 0.03a Benzaldehyde 13a Phenol 28a o-Cresol 40a 0.08a m-Cresol 57a			
p-Xylene 14a,d 0.09a, 0.08c o-Ethyltoluene 12a	•	₂₄ a,d	
o-Ethyltoluene 20°a p-Ethyltoluene 12°a 1,2,3-Trimethylbenzene 33°d 0.05°a 1,2,4-Trimethylbenzene 33°d 0.04°a 1,3,5-Trimethylbenzene 58°d 0.03°a Benzaldehyde 13°a Phenol 28°a o-Cresol 40°a 0.08°a m-Cresol 57°a	-	₁₄ 2,d	0.09 ^a , 0.08 ^c
p-Ethyltoluene 12 ^a 1,2,3-Trimethylbenzene 33 ^d 0.05 ^a 1,2,4-Trimethylbenzene 33 ^d 0.04 ^a 1,3,5-Trimethylbenzene 58 ^d 0.03 ^a Benzaldehyde 13 ^a Phenol 28 ^a o-Cresol 40 ^a 0.08 ^a m-Cresol 57 ^a	•	12 ^a	
p-Ethyltoluene 12 ^a 1,2,3-Trimethylbenzene 33 ^d 0.05 ^a 1,2,4-Trimethylbenzene 33 ^d 0.04 ^a 1,3,5-Trimethylbenzene 58 ^d 0.03 ^a Benzaldehyde 13 ^a Phenol 28 ^a o-Cresol 40 ^a 0.08 ^a m-Cresol 57 ^a	m-Ethyltoluene	20 ^a	
1,2,3-Trimethylbenzene 33 ^d 0.05 ^a 1,2,4-Trimethylbenzene 33 ^d 0.04 ^a 1,3,5-Trimethylbenzene 58 ^d 0.03 ^a Benzaldehyde 13 ^a Phenol 28 ^a o-Cresol 40 ^a 0.08 ^a m-Cresol 57 ^a	-	12 ^a	
1,3,5-Trimethylbenzene 58 ^d 0.03 ^a Benzaldehyde 13 ^a Phenol 28 ^a o-Cresol 40 ^a 0.08 ^a m-Cresol 57 ^a	1,2,3-Trimethylbenzene	33 ^d	0.05 ^a
Benzaldehyde 13 ^a Phenol 28 ^a o-Cresol 40 ^a 0.08 ^a m-Cresol 57 ^a	•	33 ^d	0.04 ^a
Phenol 28 ^a o-Cresol 40 ^a 0.08 ^a m-Cresol 57 ^a	1,3,5-Trimethylbenzene	58 ^d	0.03 ^a
o-Cresol m-Cresol 57 ^a	Benzaldehyde	13 ^a	
m-Cresol 57 ^a	Phenol	28 ^a	
m=01 C501	o-Cresol	40 ^a	0.08 ^a
p-Cresol 44ª	m-Cresol	57 ^a	
	p-Cresol	И́та	

^aAtkinson (1986a); values of $k_a/(k_a+k_b)$ estimated from the reaction kinetics. bAtkinson et al. (1983b). CBandow and Washida (1985a). dAtkinson and Aschmann, unpublished data (1988).

$$\begin{array}{c} CH_2 \\ CH_2OO \\ CH_2OO \\ CH_2OO \\ CH_2O \\ CH_2O \\ CH_2O \\ CH_2O \\ CHO \\$$

The amount of nitrate formation (benzyl nitrate from the benzyl radical) is approximately 10% (Hoshino and Akimoto, 1978; Gery et al., 1985) at room temperature and atmospheric pressure. Similar reaction pathways account for the observation of the tolualdehydes from the xylene isomers (Bandow and Washida, 1985a).

The major pathway of the OH radical reaction is hence by OH radical addition to the aromatic ring to yield hydroxycyclohexadienyl or hydroxyalkylcyclohexadienyl radicals (reaction (b) above). The hydroxycyclohexadienyl radical (HCHD) has been observed by ultraviolet absorption spectroscopy for the OH radical reaction with benzene (Fritz et al., 1985; Zellner et al., 1985) and its reactions with O_2 , NO and O_2 studied (Zellner et al., 1985). At 298 K rate constants for the reactions of HCHD radicals with NO and O_2 of

$$k(\text{HCHD} + \text{NO}) = (1.0 \pm 0.5) \times 10^{-12} \text{ cm}^3 \text{ molecule}^{-1} \text{ s}^{-1}$$
 and
$$k(\text{HCHD} + \text{NO}_2) = (8.5 \pm 2.1) \times 10^{-12} \text{ cm}^3 \text{ molecule}^{-1} \text{ s}^{-1}$$

were determined (Zellner et al., 1985). No reaction of HCHD was observed with 0_2 and, although no upper limit to the rate constant was given by Zellner et al. (1985), a value of

$$k(HCHD + O_2) < 10^{-13} cm^3 molecule^{-1} s^{-1}$$

can be estimated from their data. Since the $0_2/N0_x$ concentration ratio is typically $\gtrsim 10^6$ under atmospheric conditions, these data do not allow an assessment of whether or not the 0_2 reaction will be dominant in the atmosphere. It should be noted, however, that in a recent study Atkinson et al. (1987a) obtained data which suggest that at $N0_x$ concentrations of approximately (1-4) x 10^{14} molecule cm⁻³, the phenylhydroxycyclohexadienyl radical formed from biphenyl does not react with 0_2 .

Thus, at the present time the actual reactions removing the OH-aromatic adducts (i.e., hydroxycyclohexadienyl radicals) are not known, and it is distinctly possible that they do not react with $\rm O_2$ under polluted urban atmospheric conditions. At present the only definitive data concerning the reactions subsequent to the addition reaction of OH radicals with the aromatic hydrocarbons are the products formed. For toluene, the most studied aromatic hydrocarbon, these include o-, m- and p-cresol, m-nitrotoluene, glyoxal, methylglyoxal, PAN (a secondary product which can be formed from the reactions of methylglyoxal), $\rm CH_3COCOCH=CH_2$,

 $\label{eq:chococh=ch2} \mbox{CHOCOCH=CH$_2$, CH_3$CH=CHCH=CH$_2$, $CHOC(OH)$=CHCHO, and CH_3$COCH=CHCHO.}$

The cresol yields from toluene have been determined in three recent studies, with reported yields of 0.13 ± 0.07 (Atkinson et al., 1983b), 0.13 (Leone et al., 1985) and 0.22 (Gery et al., 1985) for o-cresol, with the m- and p-cresol yields being much lower [with the o-: m-: p- yield ratio being approximately 80:5:15 (Hoshino et al., 1978; Gery et al., 1985)]. The reaction scheme postulated to account for the cresols and m-nitrotoluene has been (Atkinson et al., 1980; Killus and Whitten, 1982)

However, the recent study of 3-nitrobiphenyl formation from biphenyl throws doubt on this scheme since both the 3-nitrobiphenyl yield and the 3-nitrobiphenyl/2-hydroxybiphenyl yield ratio were observed to be independent, within the experimental errors, of the NO_2 concentration (Atkinson et al., 1987a).

The other products whose yields have been measured with some accuracy are the α -dicarbonyls for a series of aromatic hydrocarbons, and the data are given in Table 29. In addition, the expected co-products of the α -dicarbonyls have been observed (Takagi et al., 1982; Shepson et al., 1984; Becker and Klein, 1987), but generally only in small yields.

Thus, for toluene approximately 10% of the overall reaction occurs by H atom abstraction to form, in the presence of NO, benzaldehyde and benzyl nitrate. The cresols account for a further approximately 20% of the overall OH radical reaction, and glyoxal plus methylglyoxal (plus coproducts) account for approximately 25% of the reaction, although the coproducts are not known. The remaining reaction products, accounting for approximately 45% of the overall OH radical reaction, are not quantitatively known, although a variety of ring-cleavage products have been observed (Dumdei and O'Brien, 1984; Shepson et al., 1984).

Polycyclic Aromatic Hydrocarbons (PAH)

Kinetic data are available for several of the PAH, and these data are given in Table 30. Of particular interest is that the gas-phase PAH containing two or more six-membered fused ring react with $\rm N_2O_5$ (Atkinson and Aschmann, 1987, 1988 and references therein). The OH radical (in the presence of $\rm NO_X$) and $\rm N_2O_5$ reactions lead to the formation of nitroarene products, and the yields of these products are given in Table 31.

Table 29. $\alpha ext{-Dicarbonyl Yields from the OH Radical-Initiated Reactions of a Series of Aromatic Hydrocarbons}$

	α-Dicarbonyl Yield ^a			
Aromatic	Glyoxal	Methyl- Glyoxal	Biacetyl	Reference
Benzene	0.207 ± 0.019			Tuazon et al. (1986)
Toluene	0.15 ± 0.04	0.14 ± 0.04		Bandow et al. (1985)
	0.105 ± 0.019	0.146 ± 0.006		Tuazon et al. (1986)
	0.098	0.106		Gery et al. (1985)
o-Xylene			0.18 ± 0.04	Darnall et al. (1979)
			0.260 ± 0.102	Takagi et al. (1980)
			0.137 ± 0.016	Atkinson et al. (1983b)
	0.08 ± 0.04	0.23 ± 0.03	0.10 ± 0.02	Bandow and Washida (1985a
	0.087 ± 0.012	0.246 ± 0.020		Tuazon et al. (1986)
m-Xylene	0.13 ± 0.03	0.42 ± 0.05		Bandow and Washida (1985a
	0.086 ± 0.011	0.319 ± 0.009		Tuazon et al. (1986)
p-Xylene	0.24 ± 0.02	0.12 ± 0.02		Bandow and Washida (1985a
	0.225 ± 0.039	0.105 ± 0.034		Tuazon et al. (1986)
1,2,3-Trimethyl-	0.072 ± 0.001	0.18 ± 0.01	0.45 ± 0.02	Bandow and Washida (1985b
benzene		0.152 ± 0.025	_	Tuazon et al. (1986)
1,2,4-Trimethyl-	0.078 ± 0.005	0.37 ± 0.01	0.11 ± 0.01	Bandow and Washida (1985b
benzene		0.357 ± 0.017		Tuazon et al. (1986)
1,3,5-Trimethyl-		0.64 ± 0.03		Bandow and Washida (1985b
benzene		0.602 ± 0.033		Tuazon et al. (1986)

^aIndicated error limits are two standard deviations.

Rate Constants for the Gas-Phase Reactions of PAH with OH and NO $_3$ Radicals, $\rm N_2O_5$ and $\rm O_3$ at Room Temperature Table 30.

	Rate Con	Rate Constant (cm3 molecule-1 s-1) for Reaction with	e-1 s-1) for React	tion with
РАН	НО	NO3	N205	03
Naphthalene	2.2 x 10-11 a	q	1.4 x 10-17 C	<2 x 10 ^{-19 d}
Biphenyl	$7 \times 10^{-12} a$	q	<2 x 10-19 c	$<2 \times 10^{-19} \text{ d}$
1-Methylnaphthalene	5.3 x 10 ⁻¹¹ e	ą	3.3×10^{-17} e	<1.3 x 10 ⁻¹⁹ e
2-Methylnaphthalene	$5.2 \times 10^{-11} f$	Ф	4.2 x 10-17 e	$< 4 \times 10^{-19}$ f
2,3-Dimethylnaphthalene	7.7×10^{-11} f	۵	5.7 x 10-17 e	$< 4 \times 10^{-19}$ f
Acenaphthene	1.0 x 10-10 g	4.6 x 10-13 B	5.5 x 10-17 g	<5 x 10 ^{-19 g}
Acenaphthylene	1.1 x 10-10 g	5.4 x 10-12 8	a	~5.5 x 10-16 g
Phenanthrene	$3.4 \times 10^{-11} \text{ h}$			
Anthracene	1.3 x 10 ⁻¹⁰ h			
atkinson (1986a). book reaction observed. Catkinson et al. (1987a). datkinson et al. (1984c).		eAtkin, fAtkin gAtkin hBierm	eAtkinson and Aschmann (fAtkinson and Aschmann (&Atkinson and Aschmann (hBiermann et al. (1985).	(1987). (1986). (1988).

Table 31. Nitroarene Products Formed from the Gas Phase Reactions of PAH Known to be Present in Ambient Air with OH Radicals (in the Presence of $NO_{\rm x}$) and N_2O_5 , Together with their Yields at Room Temperature and Atmospheric Pressure

	Reaction with		
PAH	ОН	N ₂ O ₅	
Naphthalene	1-Nitronaphthalene (0.3%) ^a	1-Nitronaphthalene (17%) ^a	
	2-Nitronaphthalene (0.3%) ^a	2-Nitronaphthalene (7%) ^a	
Pyrene	2-Nitropyrene (~0.5%) ^b	4-Nitropyrene (~0.3%) ^b	
	4-Nitropyrene (~0.05%) ^b		
Fluoranthene	2-Nitrofluoranthene (~5%) ^b	2-Nitrofluoranthene	
	7-Nitrofluoranthene (~0.25%) ^b	(~25-30%) ^b	
	8-Nitrofluoranthene (~0.25%) ^b		
Acephen- anthrylene	Two nitroarene isomers (~0.1%) ^{b,c}	None observed ^c	
Biphenyl	3-Nitrobiphenyl (5%) ^a	No reaction observed ^a	

aAtkinson et al. (1987a). bArey et al. (1988). cZielinska et al. (1988).

In addition (Atkinson et al., 1987a), 1- and 2-naphthol are formed from the OH radical reaction with naphthalene in approximately 5% yield each, and 2-hydroxybiphenyl is formed from biphenyl in approximately 20% yield (similar to that of o-cresol from toluene).

Aromatic Aldehydes

The only aromatic aldehyde which has been studied to any extent is benzaldehyde. The potential atmospheric reactions are photolysis and chemical reaction with OH and NO $_3$ radicals and O $_3$. No data are available concerning the O $_3$ reaction, but it is expected to be of no significance as an atmospheric loss process (Atkinson and Carter, 1984). The NO $_3$ radical reaction has a room temperature rate constant of 2.5 x 10^{-15} cm 3 molecule $^{-1}$ s $^{-1}$ (Atkinson et al., 1988b), very similar to that for acetaldehyde. This NO $_3$ radical reaction is expected to proceed by H atom abstraction from the -CHO substituent group

$$C_6H_5CHO + NO_3 + HNO_3 + C_6H_5CO$$

followed by reactions of the C_6H_5CO radical (see below).

The rate constant for the OH radical reaction at room temperature is given in Table 28. In terms of their importance as atmospheric loss processes, the OH radical reaction will dominate over the ${\rm NO_3}$ radical reaction by approximately an order of magnitude. The OH radical reaction also proceeds by H atom abstraction from the -CHO group (OH radical addition to the aromatic ring is expected to be slower than the OH radical reaction with benzene since the -CHO group is electron withdrawing). Based upon the study of the Cl atom-initiated reaction of benzaldehyde of

Niki et al. (1979), the reactions after this H atom abstraction are expected to be:

This reaction sequence is also consistent with a recent study of Tuazon et al. (unpublished results, 1988) of the NO_3 radical reaction with phenol, which yields the phenoxy radical in the initial step.

Peroxybenzoyl nitrate (PBzN) is analogous to PAN in that it is in thermal equilibrium with NO₂ and the $C_6H_5C(0)00$ radical. The thermal decomposition of PBzN has been studied by Ohta and Mizoguchi (1981) and Kenley and Hendry (1982), and rate constants of 8.5 x 10^{14} e^{-12682/T} s⁻¹ (Ohta and Mizoguchi, 1981) [2.7 x 10^{-4} s⁻¹ at 298 K] and 1.6 x 10^{15} e^{-13035/T} s⁻¹ (Kenley and Hendry, 1982) [1.6 x 10^{-4} s⁻¹ at 298 K] have been reported.

It is clear from experimental and computer modeling studies of the ${\rm NO_{X}}$ -air photooxidations of toluene that benzaldehyde photolyzes (Atkinson et al., 1980; Killus and Whitten, 1980; Gery et al., 1985; Leone et al., 1985). The absorption cross section has been given by Itoh (1987) and extends to 300 nm. The photodissociation quantum yields, the photodissociation products (in particular, whether the products are radical species or not) and their wavelength dependencies are, however, not presently known.

Phenolic Compounds

Phenol and the cresols are formed from the atmospheric degradation of benzene and toluene, respectively, and some data are available concerning the atmospheric reactions of these compounds. The potential atmospheric loss processes of phenolic compounds are reaction with NO₃ and OH radicals and with O₃, together with wet and/or dry deposition (these compounds are readily incorporated into rain and cloud water and fog). For the gasphase reactions of O₃ with the cresol isomers, room temperature rate constants of (in units of 10^{-19} cm³ molecule⁻¹ s⁻¹) 2.6, 1.9 and 4.7 for o-, m-, and p-cresol, respectively, have been measured (Atkinson and Carter, 1984). Reaction with O₃ will thus be a minor removal process for the phenolic compounds under atmospheric conditions.

The gas-phase reactions of the NO $_3$ radical with phenol and the cresols are rapid, with room temperature rate constants of: phenol, 3.6 x 10^{-12} cm 3 molecule $^{-1}$ s $^{-1}$ (Atkinson et al., 1988b); o-cresol, 2.1 x 10^{-11} cm 3 molecule $^{-1}$ s $^{-1}$ (Atkinson et al., 1984d, 1988b); m-cresol, 1.6 x 10^{-11} cm 3 molecule $^{-1}$ s $^{-1}$ (Atkinson et al., 1984d, 1988b); and p-cresol, 2.2 x 10^{-11} cm 3 molecule $^{-1}$ s $^{-1}$ (Atkinson et al., 1984d, 1988b). These reactions

appear to proceed by H atom abstraction from the O-H bonds, leading to the formation of phenoxy and methylphenoxy radicals. The subsequent reactions of these radicals have been discussed above.

The room temperature rate constants for the gas-phase reactions of the OH radical with phenol and the cresols are given in Table 28. These reactions proceed mainly by OH radical addition to the aromatic ring (Atkinson, 1986a). No detailed knowledge of the products and mechanism of these OH radical reactions is known, although pyruvic acid (CH₃COCOOH) has been reported as a product (Grosjean, 1984).

Pyruvic acid photolyzes rapidly (Grosjean, 1983, 1985) with a photolysis rate, relative to that for photolysis of NO_2 , of 0.033 under atmospheric conditions (Grosjean, 1985). The photolysis products of pyruvic acid are CO_2 and acetaldehyde (Yamamoto and Back, 1985)

$$CH_3COCOOH + hv + CO_2 + CH_3CHO$$

Clearly, further work is needed concerning the atmospheric products and mechanisms of the reactions of phenolic compounds.

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